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In Re Application of:

Nicholas C. Nicolaides, Luigi Grasso, and Philip M. Sass

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For: METHODS FOR GENERATING GENETICALLY ALTERED ANTIBODY-

PRODUCING CELL LINES WITH IMPROVED ANTIBODY

CHARACTERISTICS

☐ Provisional ☐ Design Assistant Commissioner for Patents Washington DC 20231

EXPRESS MAIL LABEL NO: EL568026574US DATE OF DEPOSIT: November 7, 2000

Sir:	
	PATENT APPLICATION TRANSMITTAL LETTER
	Transmitted herewith for filing, please find
×	A Utility Patent Application under 37 C.F.R. 1.53(b).
	It is a continuing application, as follows:
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	A Provisional Patent Application under 37 C.F.R. 1.53(c).
	A Design Patent Application (submitted in duplicate).

Includ	ing the	followi	ng:				
	Provisional Application Cover Sheet.						
×	New or Revised Specification, including pages 1_ to 41_ containing:						
	\boxtimes	Specia	lication				
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DOC	KET NO.: MOR-0003 - 3 - PATENT
	Signed Statement attached deleting inventor(s) named in the prior application.
	A Preliminary Amendment.
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	 □ A Certified Copy of each of the above applications for which priority is claimed: □ is enclosed. □ has been filed in prior application Serial No filed
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	Diskette Containing DNA/Amino Acid Sequence Information.				
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	The computer readable form in this application, is identical with that filed in Application Serial Number, filed In accordance with 37 CFR 1.821(e), please use the ☐ first-filed, ☐ last-filed or ☐ only computer readable form filed in that application as the computer readable form for the instant application. It is understood that the Patent and Trademark Office will make the necessary change in application number and filing date for the computer readable form that will be used for the instant application. A paper copy of the Sequence Listing is ☐ included in the originally-filed specification of the instant application, ☐ included in a separately filed preliminary amendment for incorporation into the specification.				
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Date: 11 /7 /00

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METHODS FOR GENERATING GENETICALLY ALTERED ANTIBODY-PRODUCING CELL LINES WITH IMPROVED ANTIBODY CHARACTERISTICS

TECHNICAL FIELD OF THE INVENTION

The invention is related to the area of antibody maturation and cellular production. In particular, it is related to the field of mutagenesis.

BACKGROUND OF THE INVENTION

The use of antibodies to block the activity of foreign and/or endogenous polypeptides provides an effective and selective strategy for treating the underlying cause of disease. In particular is the use of monoclonal antibodies (MAb) as effective therapeutics such as the FDA approved ReoPro (Glaser, V. (1996) Can ReoPro repolish tarnished monoclonal therapeutics? *Nat. Biotechnol.* 14:1216-1217), an anti-platelet MAb from Centocor; Herceptin (Weiner, L.M. (1999) Monoclonal antibody therapy of cancer. *Semin. Oncol.* 26:43-51), an anti-Her2/neu MAb from Genentech; and Synagis (Saez-Llorens, X.E., et al. (1998) Safety and pharmacokinetics of an intramuscular humanized monoclonal antibody to respiratory syncytial virus in premature infants and infants with bronchopulmonary dysplasia. *Pediat. Infect. Dis. J.* 17:787-791), an anti-respiratory syncytial virus MAb produced by Medimmune.

Standard methods for generating MAbs against candidate protein targets are known by those skilled in the art. Briefly, rodents such as mice or rats are injected with a purified antigen in the presence of adjuvant to generate an immune response (Shield, C.F., et al. (1996) A cost-effective analysis of OKT3 induction therapy in cadaveric kidney

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transplantation. Am. J. Kidney Dis. 27:855-864). Rodents with positive immune sera are sacrificed and splenocytes are isolated. Isolated splenocytes are fused to melanomas to produce immortalized cell lines that are then screened for antibody production. Positive lines are isolated and characterized for antibody production. The direct use of rodent MAbs as human therapeutic agents were confounded by the fact that human anti-rodent antibody (HARA) responses occurred in a significant number of patients treated with the rodentderived antibody (Khazaeli, M.B., et al., (1994) Human immune response to monoclonal antibodies. J. Immunother. 15:42-52). In order to circumvent the problem of HARA, the grafting of the complementarity determining regions (CDRs), which are the critical motifs found within the heavy and light chain variable regions of the immunoglobulin (Ig) subunits making up the antigen binding domain, onto a human antibody backbone found these chimeric molecules are able to retain their binding activity to antigen while lacking the HARA response (Emery, S.C., and Harris, W.J. "Strategies for humanizing antibodies" In: ANTIBODY ENGINEERING C.A.K. Borrebaeck (Ed.) Oxford University Press, N.Y. 1995. pp. 159-183. A common problem that exists during the "humanization" of rodent-derived MAbs (referred to hereon as HAb) is the loss of binding affinity due to conformational changes in the 3 dimensional structure of the CDR domain upon grafting onto the human Ig backbone (U.S. Patent No. 5,530,101 to Queen et al.). To overcome this problem, additional HAb vectors are usually needed to be engineered by inserting or deleting additional amino acid residues within the framework region and/or within the CDR coding region itself in order to recreate high affinity HAbs (U.S. Patent No. 5,530,101 to Queen et al.). This process is a very time consuming procedure that involves the use of expensive computer modeling programs to predict changes that may lead to a high affinity HAb. In some instances the affinity of the HAb is never restored to that of the MAb, rendering them of little therapeutic use.

Another problem that exists in antibody engineering is the generation of stable, high yielding producer cell lines that is required for manufacturing of the molecule for clinical materials. Several strategies have been adopted in standard practice by those skilled in the art to circumvent this problem. One method is the use of Chinese Hamster Ovary (CHO) cells transfected with exogenous Ig fusion genes containing the grafted human light and heavy

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chains to produce whole antibodies or single chain antibodies, which are a chimeric molecule containing both light and heavy chains that form an antigen-binding polypeptide (Reff, M.E. (1993) High-level production of recombinant immunoglobulins in mammalian cells. *Curr. Opin. Biotechnol.* 4:573-576). Another method employs the use of human lymphocytes derived from transgenic mice containing a human grafted immune system or transgenic mice containing a human Ig gene repertoire. Yet another method employs the use of monkeys to produce primate MAbs, which have been reported to lack a human anti-monkey response (Neuberger, M., and Gruggermann, M. (1997) Monoclonal antibodies. Mice perform a human repertoire. *Nature* 386:25-26). In all cases, the generation of a cell line that is capable of generating sufficient amounts of high affinity antibody poses a major limitation for producing sufficient materials for clinical studies. Because of these limitations, the utility of other recombinant systems such as plants are currently being explored as systems that will lead to the stable, high-level production of humanized antibodies (Fiedler, U., and Conrad, U. (1995) High-level production and long-term storage of engineered antibodies in transgenic tobacco seeds. *Bio/Technology* 13:1090-1093).

A method for generating diverse antibody sequences within the variable domain that results in HAbs and MAbs with high binding affinities to antigens would be useful for the creation of more potent therapeutic and diagnostic reagents respectively. Moreover, the generation of randomly altered nucleotide and polypeptide residues throughout an entire antibody molecule will result in new reagents that are less antigenic and/or have beneficial pharmacokinetic properties. The invention described herein is directed to the use of random genetic mutation throughout an antibody structure in vivo by blocking the endogenous mismatch repair (MMR) activity of a host cell producing immunoglobulins that encode biochemically active antibodies. The invention also relates to methods for repeated in vivo genetic alterations and selection for antibodies with enhanced binding and pharmacokinetic profiles.

In addition, the ability to develop genetically altered host cells that are capable of secreting increased amounts of antibody will also provide a valuable method for creating cell hosts for product development. The invention described herein is directed to the creation of genetically altered cell hosts with increased antibody production via the blockade of MMR.

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The invention facilitates the generation of high affinity antibodies and the production of cell lines with elevated levels of antibody production. Other advantages of the present invention are described in the examples and figures described herein.

5 SUMMARY OF THE INVENTION

The invention provides methods for generating genetically altered antibodies (including single chain molecules) and antibody producing cell hosts in vitro and in vivo, whereby the antibody possess a desired biochemical property(s), such as, but not limited to, increased antigen binding, increased gene expression, and/or enhanced extracellular secretion by the cell host. One method for identifying antibodies with increased binding activity or cells with increased antibody production is through the screening of MMR defective antibody producing cell clones that produce molecules with enhanced binding properties or clones that have been genetically altered to produce enhanced amounts of antibody product.

The antibody producing cells suitable for use in the invention include, but are not limited to rodent, primate, or human hybridomas or lymphoblastoids; mammalian cells transfected and expressing exogenous Ig subunits or chimeric single chain molecules; plant cells, yeast or bacteria transfected and expressing exogenous Ig subunits or chimeric single chain molecules.

Thus, the invention provides methods for making hypermutable antibody-producing cells by introducing a polynucleotide comprising a dominant negative allele of a mismatch repair gene into cells that are capable of producing antibodies. The cells that are capable of producing antibodies include cells that naturally produce antibodies, and cells that are engineered to produce antibodies through the introduction of immunoglobulin encoding sequences. Conveniently, the introduction of polynucleotide sequences into cells is accomplished by transfection.

The invention also provides methods of making hypermutable antibody producing cells by introducing a dominant negative mismatch repair (MMR) gene such as *PMS2* (preferably human *PMS2*), *MLH1*, *PMS1*, *MSH2*, or *MSH2* into cells that are capable of producing antibodies. The dominant negative allele of a mismatch repair gene may be a truncation mutation of a mismatch repair gene (preferably a truncation mutation at codon 134,

or a thymidine at nucleotide 424 of wild-type *PMS2*). The invention also provides methods in which mismatch repair gene activity is suppressed. This may be accomplished, for example, using antisense molecules directed against the mismatch repair gene or transcripts.

Other embodiments of the invention provide methods for making a hypermutable antibody producing cells by introducing a polynucleotide comprising a dominant negative allele of a mismatch repair gene into fertilized eggs of animals. These methods may also include subsequently implanting the eggs into pseudo-pregnant females whereby the fertilized eggs develop into a mature transgenic animal. The mismatch repair genes may include, for example, PMS2 (preferably human PMS2), MLH1, PMS1, MSH2, or MSH2. The dominant negative allele of a mismatch repair gene may be a truncation mutation of a mismatch repair gene (preferably a truncation mutation at codon 134, or a thymidine at nucleotide 424 of wild-type PMS2).

The invention further provides homogeneous compositions of cultured, hypermutable, mammalian cells that are capable of producing antibodies and contain a dominant negative allele of a mismatch repair gene. The mismatch repair genes may include, for example, *PMS2* (preferably human *PMS2*), *MLH1*, *PMS1*, *MSH2*, or *MSH2*. The dominant negative allele of a mismatch repair gene may be a truncation mutation of a mismatch repair gene (preferably a truncation mutation at codon 134, or a thymidine at nucleotide 424 of wild-type *PMS2*). The cells of the culture may contain *PMS2*, (preferably human *PMS2*), *MLH1*, or *PMS1*; or express a human *mutL* homolog, or the first 133 amino acids of hPMS2.

The invention further provides methods for generating a mutation in an immunoglobulin gene of interest by culturing an immunoglobulin producing cell selected for an immunoglobulin of interest wherein the cell contains a dominant negative allele of a mismatch repair gene. The properties of the immunoglobulin produced from the cells can be assayed to ascertain whether the immunoglobulin gene harbors a mutation. The assay may be directed to analyzing a polynucleotide encoding the immunoglobulin, or may be directed to the immunoglobulin polypeptide itself.

The invention also provides methods for generating a mutation in a gene affecting antibody production in an antibody-producing cell by culturing the cell expressing a dominant negative allele of a mismatch repair gene, and testing the cell to determine whether the cell

harbors mutations within the gene of interest, such that a new biochemical feature (e.g., overexpression and/or secretion of immunoglobulin products) is generated. The testing may include analysis of the steady state expression of the immunoglobulin gene of interest, and/or analysis of the amount of secreted protein encoded by the immunoglobulin gene of interest. The invention also embraces prokaryotic and eukaryotic transgenic cells made by this process, including cells from rodents, non-human primates and humans.

Other aspects of the invention encompass methods of reversibly altering the hypermutability of an antibody producing cell, in which an inducible vector containing a dominant negative allele of a mismatch repair gene operably linked to an inducible promoter is introduced into an antibody-producing cell. The cell is treated with an inducing agent to express the dominant negative mismatch repair gene (which can be *PMS2* (preferably human *PMS2*), *MLH1*, or *PMS1*). Alternatively, the cell may be induced to express a human *mutL* homolog or the first 133 amino acids of hPMS2. In another embodiment, the cells may be rendered capable of producing antibodies by co-transfecting a preselected immunoglobulin gene of interest. The immunoglobulin genes of the hypermutable cells, or the proteins produced by these methods may be analyzed for desired properties, and induction may be stopped such that the genetic stability of the host cell is restored.

The invention also embraces methods of producing genetically altered antibodies by transfecting a polynucleotide encoding an immunoglobulin protein into a cell containing a dominant negative mismatch repair gene (either naturally or in which the dominant negative mismatch repair gene was introduced into the cell), culturing the cell to allow the immunoglobulin gene to become mutated and produce a mutant immunoglobulin, screening for a desirable property of said mutant immunoglobulin protein, isolating the polynucleotide molecule encoding the selected mutant immunoglobulin possessing the desired property, and transfecting said mutant polynucleotide into a genetically stable cell, such that the mutant antibody is consistently produced without further genetic alteration. The dominant negative mismatch repair gene may be *PMS2* (preferably human *PMS2*), *MLH1*, or *PMS1*.

Alternatively, the cell may express a human *mutL* homolog or the first 133 amino acids of hPMS2.

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The invention further provides methods for generating genetically altered cell lines that express enhanced amounts of an antigen binding polypeptide. These antigen-binding polypeptides may be, for example, immunoglobulins. The methods of the invention also include methods for generating genetically altered cell lines that secrete enhanced amounts of an antigen binding polypeptide. The cell lines are rendered hypermutable by dominant negative mismatch repair genes that provide an enhanced rate of genetic hypermutation in a cell producing antigen-binding polypeptides such as antibodies. Such cells include, but are not limited to hybridomas. Expression of enhanced amounts of antigen binding polypeptides may be through enhanced transcription or translation of the polynucleotides encoding the antigen binding polypeptides, or through the enhanced secretion of the antigen binding polypeptides, for example.

Methods are also provided for creating genetically altered antibodies in vivo by blocking the MMR activity of the cell host, or by transfecting genes encoding for immunoglobulin in a MMR defective cell host.

Antibodies with increased binding properties to an antigen due to genetic changes within the variable domain are provided in methods of the invention that block endogenous MMR of the cell host. Antibodies with increased binding properties to an antigen due to genetic changes within the CDR regions within the light and/or heavy chains are also provided in methods of the invention that block endogenous MMR of the cell host.

The invention provides methods of creating genetically altered antibodies in MMR defective Ab producer cell lines with enhanced pharmacokinetic properties in host organisms including but not limited to rodents, primates, and man.

These and other aspects of the invention are provided by one or more of the embodiments described below. In one embodiment of the invention, a method for making an antibody producing cell line hypermutable is provided. A polynucleotide encoding a dominant negative allele of a MMR gene is introduced into an antibody-producing cell. The cell becomes hypermutable as a result of the introduction of the gene.

In another embodiment of the invention, a method is provided for introducing a mutation into an endogenous gene encoding for an immunoglobulin polypeptide or a single chain antibody. A polynucleotide encoding a dominant negative allele of a MMR gene is

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introduced into a cell. The cell becomes hypermutable as a result of the introduction and expression of the MMR gene allele. The cell further comprises an immunoglobulin gene of interest. The cell is grown and tested to determine whether the gene encoding for an immunoglobulin or a single chain antibody of interest harbors a mutation. In another aspect of the invention, the gene encoding the mutated immunoglobulin polypeptide or single chain antibody may be isolated and expressed in a genetically stable cell. In a preferred embodiment, the mutated antibody is screened for at least one desirable property such as, but not limited to, enhanced binding characteristics.

In another embodiment of the invention, a gene or set of genes encoding for Ig light and heavy chains or a combination therein are introduced into a mammalian cell host that is MMR defective. The cell is grown, and clones are analyzed for antibodies with enhanced binding characteristics.

In another embodiment of the invention, a method will be provided for producing new phenotypes of a cell. A polynucleotide encoding a dominant negative allele of a MMR gene is introduced into a cell. The cell becomes hypermutable as a result of the introduction of the gene. The cell is grown. The cell is tested for the expression of new phenotypes where the phenotype is enhanced secretion of a polypeptide.

These and other embodiments of the invention provide the art with methods that can generate enhanced mutability in cells and animals as well as providing cells and animals harboring potentially useful mutations for the large-scale production of high affinity antibodies with beneficial pharmacokinetic profiles.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1. Hybridoma cells stably expressing PMS2 and PMS134 MMR genes.

Shown is steady state mRNA expression of MMR genes transfected into a murine hybridoma cell line. Stable expression was found after 3 months of continuous growth. The (-) lanes represent negative controls where no reverse transcriptase was added, and the (+) lanes represent samples reverse transcribed and PCR amplified for the MMR genes and an internal housekeeping gene as a control.

Figure 2. Creation of genetically hypermutable hybridoma cells. Dominant negative

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MMR gene alleles were expressed in cells expressing a MMR-sensitive reporter gene. Dominant negative alleles such as PMS134 and the expression of MMR genes from other species results in antibody producer cells with a hypermutable phenotype that can be used to produce genetically altered immunoglobulin genes with enhanced biochemical features as well as lines with increased Ig expression and/or secretion. Values shown represent the amount of converted CPRG substrate which is reflective of the amount of function \Box -galactosidase contained within the cell from genetic alterations within the pCAR-OF reporter gene. Higher amounts of β -galactosidase activity reflect a higher mutation rate due to defective MMR.

Figure 3. Screening method for identifying antibody-producing cells containing antibodies with increased binding activity and/or increased expression/secretion

Figure 4. Generation of a genetically altered antibody with an increased binding activity. Shown are ELISA values from 96-well plates, screened for antibodies specific to hIgE. Two clones with a high binding value were found in HB134 cultures.

Figure 5. Sequence alteration within variable chain of an antibody (a mutation within the light chain variable region in MMR-defective HB134 antibody producer cells). Arrows indicate the nucleotide at which a mutation occurred in a subset of cells from a clone derived from HB134 cells. Panel A: The change results in a Thr to Ser change within the light chain variable region. The coding sequence is in the antisense direction. Panel B: The change results in a Pro to His change within the light chain variable region.

Figure 6. Generation of MMR-defective clones with enhanced steady state Ig protein levels. A Western blot of heavy chain immunglobulins from HB134 clones with high levels of MAb (>500ngs/ml) within the conditioned medium shows that a subset of clones express higher steady state levels of immunoglobulins (Ig). The H36 cell line was used as a control to measure steady state levels in the parental strain. Lane 1: fibroblast cells (negative control); Lane 2: H36 cell; Lane 3: HB134 clone with elevated MAb levels; Lane 4: HB134 clone with elevated MAb levels.

Methods have been discovered for developing hypermutable antibody-producing cells by taking advantage of the conserved mismatch repair (MMR) process of host cells.

30 Dominant negative alleles of such genes, when introduced into cells or transgenic animals,

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increase the rate of spontaneous mutations by reducing the effectiveness of DNA repair and thereby render the cells or animals hypermutable. Hypermutable cells or animals can then be utilized to develop new mutations in a gene of interest. Blocking MMR in antibodyproducing cells such as but not limited to; hybridomas; mammalian cells transfected with genes encoding for Ig light and heavy chains; mammalian cells transfected with genes encoding for single chain antibodies; eukaryotic cells transfected with Ig genes, can enhance the rate of mutation within these cells leading to clones that have enhanced antibody production and/or cells containing genetically altered antibodies with enhanced biochemical properties such as increased antigen binding. The process of MMR, also called mismatch proofreading, is carried out by protein complexes in cells ranging from bacteria to mammalian cells. A MMR gene is a gene that encodes for one of the proteins of such a mismatch repair complex. Although not wanting to be bound by any particular theory of mechanism of action, a MMR complex is believed to detect distortions of the DNA helix resulting from noncomplementary pairing of nucleotide bases. The non-complementary base on the newer DNA strand is excised, and the excised base is replaced with the appropriate base, which is complementary to the older DNA strand. In this way, cells eliminate many mutations that occur as a result of mistakes in DNA replication.

Dominant negative alleles cause a MMR defective phenotype even in the presence of a wild-type allele in the same cell. An example of a dominant negative allele of a MMR gene is the human gene hPMS2-134, which carries a truncating mutation at codon 134 (SEQ ID NO:15). The mutation causes the product of this gene to abnormally terminate at the position of the 134th amino acid, resulting in a shortened polypeptide containing the N-terminal 133 amino acids. Such a mutation causes an increase in the rate of mutations, which accumulate in cells after DNA replication. Expression of a dominant negative allele of a mismatch repair gene results in impairment of mismatch repair activity, even in the presence of the wild-type allele. Any allele which produces such effect can be used in this invention. Dominant negative alleles of a MMR gene can be obtained from the cells of humans, animals, yeast, bacteria, or other organisms. Such alleles can be identified by screening cells for defective MMR activity. Cells from animals or humans with cancer can be screened for defective mismatch repair. Cells from colon cancer patients may be particularly useful. Genomic

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DNA, cDNA, or mRNA from any cell encoding a MMR protein can be analyzed for variations from the wild type sequence. Dominant negative alleles of a MMR gene can also be created artificially, for example, by producing variants of the hPMS2-134 allele or other MMR genes. Various techniques of site-directed mutagenesis can be used. The suitability of such alleles, whether natural or artificial, for use in generating hypermutable cells or animals can be evaluated by testing the mismatch repair activity caused by the allele in the presence of one or more wild-type alleles, to determine if it is a dominant negative allele.

A cell or an animal into which a dominant negative allele of a mismatch repair gene has been introduced will become hypermutable. This means that the spontaneous mutation rate of such cells or animals is elevated compared to cells or animals without such alleles. The degree of elevation of the spontaneous mutation rate can be at least 2-fold, 5-fold, 10-fold, 20-fold, 50-fold, 50-fold, 50-fold, or 1000-fold that of the normal cell or animal. The use of chemical mutagens such as but limited to methane sulfonate, dimethyl sulfonate, O6-methyl benzadine, MNU, ENU, etc. can be used in MMR defective cells to increase the rates an additional 10 to 100 fold that of the MMR deficiency itself.

According to one aspect of the invention, a polynucleotide encoding for a dominant negative form of a MMR protein is introduced into a cell. The gene can be any dominant negative allele encoding a protein, which is part of a MMR complex, for example, *PMS2*, *PMS1*, *MLH1*, or *MSH2*. The dominant negative allele can be naturally occurring or made in the laboratory. The polynucleotide can be in the form of genomic DNA, cDNA, RNA, or a chemically synthesized polynucleotide.

The polynucleotide can be cloned into an expression vector containing a constitutively active promoter segment (such as but not limited to CMV, SV40, Elongation Factor or LTR sequences) or to inducible promoter sequences such as the steroid inducible pIND vector (Invitrogen), where the expression of the dominant negative MMR gene can be regulated. The polynucleotide can be introduced into the cell by transfection.

According to another aspect of the invention, an immunoglobulin (Ig) gene, a set of Ig genes or a chimeric gene containing whole or parts of an Ig gene can be transfected into MMR deficient cell hosts, the cell is grown and screened for clones containing genetically altered Ig genes with new biochemical features. MMR defective cells may be of human,

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primates, mammals, rodent, plant, yeast or of the prokaryotic kingdom. The mutated gene encoding the Ig with new biochemical features may be isolated from the respective clones and introduced into genetically stable cells (i.e., cells with normal MMR) to provide clones that consistently produce Ig with the new biochemical features. The method of isolating the Ig gene encoding Ig with new biochemical features may be any method known in the art. Introduction of the isolated polynucleotide encoding the Ig with new biochemical features may also be performed using any method known in the art, including, but not limited to transfection of an expression vector containing the polynucleotide encoding the Ig with new biochemical features. As an alternative to transfecting an Ig gene, a set of Ig genes or a chimeric gene containing whole or parts of an Ig gene into an MMR deficient host cell, such Ig genes may be transfected simultaneously with a gene encoding a dominant negative mismatch repair gene into a genetically stable cell to render the cell hypermutable.

Transfection is any process whereby a polynucleotide is introduced into a cell. The process of transfection can be carried out in a living animal, e.g., using a vector for gene therapy, or it can be carried out in vitro, e.g., using a suspension of one or more isolated cells in culture. The cell can be any type of eukaryotic cell, including, for example, cells isolated from humans or other primates, mammals or other vertebrates, invertebrates, and single celled organisms such as protozoa, yeast, or bacteria.

In general, transfection will be carried out using a suspension of cells, or a single cell, but other methods can also be applied as long as a sufficient fraction of the treated cells or tissue incorporates the polynucleotide so as to allow transfected cells to be grown and utilized. The protein product of the polynucleotide may be transiently or stably expressed in the cell. Techniques for transfection are well known. Available techniques for introducing polynucleotides include but are not limited to electroporation, transduction, cell fusion, the use of calcium chloride, and packaging of the polynucleotide together with lipid for fusion with the cells of interest. Once a cell has been transfected with the MMR gene, the cell can be grown and reproduced in culture. If the transfection is stable, such that the gene is expressed at a consistent level for many cell generations, then a cell line results.

An isolated cell is a cell obtained from a tissue of humans or animals by mechanically separating out individual cells and transferring them to a suitable cell culture medium, either

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with or without pretreatment of the tissue with enzymes, e.g., collagenase or trypsin. Such isolated cells are typically cultured in the absence of other types of cells. Cells selected for the introduction of a dominant negative allele of a mismatch repair gene may be derived from a eukaryotic organism in the form of a primary cell culture or an immortalized cell line, or may be derived from suspensions of single-celled organisms.

A polynucleotide encoding for a dominant negative form of a MMR protein can be introduced into the genome of an animal by producing a transgenic animal. The animal can be any species for which suitable techniques are available to produce transgenic animals. For example, transgenic animals can be prepared from domestic livestock, e.g., bovine, swine, sheep, goats, horses, etc.; from animals used for the production of recombinant proteins, e.g., bovine, swine, or goats that express a recombinant polypeptide in their milk; or experimental animals for research or product testing, e.g., mice, rats, guinea pigs, hamsters, rabbits, etc. Cell lines that are determined to be MMR defective can then be used as a source for producing genetically altered immunoglobulin genes in vitro by introducing whole, intact immunoglobulin genes and/or chimeric genes encoding for single chain antibodies into MMR defective cells from any tissue of the MMR defective animal.

Once a transfected cell line or a colony of transgenic animals has been produced, it can be used to generate new mutations in one or more gene(s) of interest. A gene of interest can be any gene naturally possessed by the cell line or transgenic animal or introduced into the cell line or transgenic animal. An advantage of using such cells or animals to induce mutations is that the cell or animal need not be exposed to mutagenic chemicals or radiation, which may have secondary harmful effects, both on the object of the exposure and on the workers. However, chemical mutagens may be used in combination with MMR deficiency, which renders such mutagens less toxic due to an undetermined mechanism. Hypermutable animals can then be bred and selected for those producing genetically variable B-cells that may be isolated and cloned to identify new cell lines that are useful for producing genetically variable cells. Once a new trait is identified, the dominant negative MMR gene allele can be removed by directly knocking out the allele by technologies used by those skilled in the art or by breeding to mates lacking the dominant negative allele to select for offspring with a desired trait and a stable genome. Another alternative is to use a CRE-LOX expression

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system, whereby the dominant negative allele is spliced from the animal genome once an animal containing a genetically diverse immunoglobulin profile has been established. Yet another alternative is the use of inducible vectors such as the steroid induced pIND (Invitrogen) or pMAM (Clonetech) vectors which express exogenous genes in the presence of corticosteroids.

Mutations can be detected by analyzing for alterations in the genotype of the cells or animals, for example by examining the sequence of genomic DNA, cDNA, messenger RNA, or amino acids associated with the gene of interest. Mutations can also be detected by screening for the production of antibody titers. A mutant polypeptide can be detected by identifying alterations in electrophoretic mobility, spectroscopic properties, or other physical or structural characteristics of a protein encoded by a mutant gene. One can also screen for altered function of the protein in situ, in isolated form, or in model systems. One can screen for alteration of any property of the cell or animal associated with the function of the gene of interest, such as but not limited to Ig secretion.

Examples of mismatch repair proteins and nucleic acid sequences include the following:

PMS2 (mouse) (SEQ ID NO:5)

```
MEOTEGVSTE CAKAIKPIDG KSVHOICSGQ VILSLSTAVK ELIENSVDAG ATTIDLRLKD
    YGVDLIEVSD NGCGVEEENF EGLALKHHTS KIOEFADLTO VETFGFRGEA LSSLCALSDV 120
20
    TISTCHGSAS VGTRLVFDHN GKITOKTPYP RPKGTTVSVO HLFYTLPVRY KEFORNIKKE 180
    YSKMVQVLQA YCIISAGVRV SCTNQLGQGK RHAVVCTSGT SGMKENIGSV FGQKQLQSLI 240
     PFVQLPPSDA VCEEYGLSTS GRHKTFSTFR ASFHSARTAP GGVQQTGSFS SSIRGPVTQQ 300
    RSLSLSMRFY HMYNRHQYPF VVLNVSVDSE CVDINVTPDK RQILLQEEKL LLAVLKTSLI 360
25
    GMFDSDANKL NVNOOPLLDV EGNLVKLHTA ELEKPVPGKO DNSPSLKSTA DEKRVASISR 420
     LREAFSLHPT KEIKSRGPET AELTRSFPSE KRGVLSSYPS DVISYRGLRG SODKLVSPTD 480
     SPGDCMDREK IEKDSGLSST SAGSEEEFST PEVASSFSSD YNVSSLEDRP SOETINCGDL 540
     DCRPPGTGOS LKPEDHGYOC KALPLARLSP TNAKRFKTEE RPSNVNISOR LPGPOSTSAA 600
     EVDVAIKMNK RIVLLEFSLS SLAKRMKQLQ HLKAQNKHEL SYRKFRAKIC PGENQAAEDE 660
30
    LRKEISKSMF AEMEILGOFN LGFIVTKLKE DLFLVDQHAA DEKYNFEMLQ QHTVLQAQRL 720
     ITPOTLNLTA VNEAVLIENL EIFRKNGFDF VIDEDAPVTE RAKLISLPTS KNWTFGPODI 780
     DELIFMLSDS PGVMCRPSRV ROMFASRACR KSVMIGTALN ASEMKKLITH MGEMDHPWNC 840
     PHGRPTMRHV ANLDVISQN
                                                                       859
```

35 PMS2 (mouse cDNA) (SEO ID NO:6)

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	gtcttttccc	gagageggea	ccgcaactct	cccgcggtga	ctgtgactgg	aggagtcctg	180
40	catccatgga	gcaaaccgaa	ggcgtgagta	cagaatqtgc	taaggccatc	aagcctattg	240
	atgggaagtc	agtccatcaa	atttgttctg	ggcaggtgat	actcagttta	agcaccgctg	300
	tgaaggagtt	gatagaaaat	agtgtagatg	ctggtgctac	tactattgat	ctaaggctta	360
	aagactatgg	ggtggacctc	attgaagttt	cagacaatgg	atgtggggta	gaagaagaaa	420

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actttgaagg totagototg aaacatcaca catotaagat toaagagttt googacotca 480
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     ataatqqqaa aatcacccaq aaaactccct acccccqacc taaaqqaacc acaqtcaqtq 660
     tgcagcactt attttataca ctacccgtgc gttacaaaga gtttcagagg aacattaaaa 720
     aggadtatte caaaatggtg caggtettac aggegtactg tateatetea geaggegtee 780
     qtqtaaqctq cactaatcaq ctcggacagg ggaagcggca cgctgtggtg tgcacaagcg 840
     gcacqtctqq catqaaqqaa aatatcqqqt ctqtqtttqq ccaqaaqcaq ttqcaaaqcc 900
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     cgccqqqaqq agtqcaacaq acaggcagtt tttcttcatc aatcagaggc cctgtgaccc 1080
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     cattigting cottaacqtt toogttgact cagaatqtgt ggatattaat gtaactocag 1200
     ataaaaggca aattotacta caagaagaga agotattgot ggccgtttta aagacctcct 1260
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     atqttqaaqq taacttaqta aagctgcata ctgcagaact agaaaagcct qtqccaqqaa 1380
     agcaagataa ctctccttca ctgaagagca cagcagacga gaaaagggta gcatccatct 1440
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     agactgctga actgacacgg agttttccaa gtgagaaaag gggcgtgtta tcctcttatc 1560
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     aactgagtta cagaaaattt agggccaaga tttgccctgg agaaaaccaa gcagcagaag 2160
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     ctgcqqatqa qaaqtacaac tttgagatgc tgcagcagca cacggtgctc caggcgcaga 2340
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     PMS2 (human) (SEQ ID NO:7)
     MERAESSSTE PAKAIKPIDR KSVHQICSGQ VVLSLSTAVK ELVENSLDAG ATNIDLKLKD 60
     YGVDLIEVSD NGCGVEEENF EGLTLKHHTS KIQEFADLTQ VETFGFRGEA LSSLCALSDV 120
     TISTCHASAK VGTRLMFDHN GKIIQKTPYP RPRGTTVSVQ QLFSTLPVRH KEFQRNIKKE 180
YAKMVQVLHA YCIISAGIRV SCTNQLGQGK RQPVVCTGGS PSIKENIGSV FGQKQLQSLI 240
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     PFVQLPPSDS VCEEYGLSCS DALHNLFYIS GFISQCTHGV GRSSTDRQFF FINRRPCDPA 300
     KYCRI, VNEVY HMYNRHOYPF VYLNISVDSE CVDINYTPDK ROILLOEEKL LLAVLKTSLI 360
     GMFDSDVNKL NVSQQPLLDV EGNLIKMHAA DLEKPMVEKQ DQSPSLRTGE EKKDVSISRL 420
     REAFSLRHTT ENKPHSPKTP EPRRSPLGQK RGMLSSSTSG AISDKGVLRP QKEAVSSSHG 480
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     PSDPTDRAEV EKDSGHGSTS VDSEGFSIPD TGSHCSSEYA ASSPGDRGSQ EHVDSQEKAP 540
     ETDDSFSDVD CHSNQEDTGC KFRVLPQPTN LATPNTKRFK KEELLSSDI CQKLVNTQDM 600
SASQVDVAVK INKKVVPLDF SMSSLAKRIK QLHHBAQQSE GBQNYRKFRA KICPGENQAA 660
EDBLRKEISK TMFARMEIIG QFNLGFLITTK LMEDIFIVDQ HATDBKYNFF MLQQHTVLQG 720
     ORLIAPOTLN LTAVNEAVLI ENLEIFRKNG FDFVIDENAP VTERAKLISL PTSKNWTFGP 780
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     WNCPHGRPTM RHIANLGVIS QN
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PMS2 (human cDNA) (SEO ID NO:8)
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     ctgagtctaa gcactgcggt aaaggagtta gtagaaaaca gtctggatgc tggtgccact 180
     aatattgato taaagottaa ggactatgga gtggatotta ttgaagttto aqacaatqqa 240
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     tttatcgcag atttttatgt tttgaaagac agagtcttca ctaacctttt ttgttttaaa 2700
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     cttttcaaac c
50
     PMS1 (human) (SEQ ID NO:9)
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     IKAVDAPVMA MKYYTSKINS HEDLENLTTY GFRGEALGSI CCIAEVLITT RTAADNFSTQ 120
YVLDGSGHIL SQKPSHLGQG TTVTALRLFK NLPVRKQFYS TAKKCKDEIK KIQDLLMSFG 180
     ILKPDLRIVF VHNKAVIWOK SRVSDHKMAL MSVLGTAVMN NMESFQYHSE ESQIYLSGFL 240
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     PKCDADHSFT SLSTPERSFI FINSRPVHOK DILKLIRHHY NLKCLKESTR LYPVFFLKID 300
     VPTADVDVNL TPDKSOVLLO NKESVLIALE NLMTTCYGPL PSTNSYENNK TDVSAADIVL 360
     SKTAETDVLF NKVESSGKNY SNVDTSVIPF QNDMHNDESG KNTDDCLNHQ ISIGDFGYGH 420
CSSEISNIDK NTKNAFQDIS MSNVSWENSQ TEYSKTCFIS SVKHTQSENG NKDHIDESGE 480
60
     NEEEAGLENS SEISADEWSR GNILKNSVGE NIEPVKILVP EKSLPCKVSN NNYPIPEQMN 540
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	NLAQKHKLKT KDEPCLIHNL	SLSNQPKLDE RFPDAWLMTS	LLQSQIEKRR KTEVMLLNPY	SQNIKMVQIP RVEEALLFKR	FSMKNLKINF LLENHKLPAE		720 780
5		KEILNAILNR EIKECVHGRP			AVRLSRQLPM	YLSKEDIQDI	900 932
	PMS1 (huma	n) (SEQ ID N	VO:10)				
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10						ctcctttcaa	
						aactccttgg	
						aaaattgagg	
						atgaagtact	
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						aatctacctq	
						aagatccaag	
						gtacataaca	
20						atgtcagttc	
						gaatctcaga	
	tttatctcag	tggatttctt	ccaaagtgtg	atgcagacca	ctctttcact	agtctttcaa	840
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						acagatgttt	
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						aacactaaga	
						acggaatata	
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35						aatattgaac	
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						aatgtaatag	
						atcaagaaac	
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50						cctcgtctta	
						aaagaaattc	
						cgcaaagtga	
						tacttatcaa	
55						gaaattaaag	
						actacatgat	
						acagcatgag	
						tgaggattca	
60		tttatattga	aaaaagttcc	acgtattgta	gaaaacgtaa	ataaactaat	
00	aac						3063

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MSH2 (human) (SEQ ID NO:11)

5							
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						RKLGLCEFPD	
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						LTTFDFSQYM	
10						MDKNRIEERL	
						INQLPNVIQA	
						KPSFDPNLSE	
						KVLRNNKNFS	
						PMQTLNDVLA	
15	QLDAVVSFAH	VSNGAPVPYV	RPAILEKGQG	RIILKASRHA	CVEVQDEIAF	IPNDVYFEKD	660
	KQMFHIITGP	NMGGKSTYIR	QTGVIVLMAQ	IGCFVPCESA	EVSIVDCILA	RVGAGDSQLK	720
						ATKIGAFCMF	
						AELANFPKHV	
	IECAKQKALE	LEEFQYIGES	QGYDIMEPAA	KKCYLEREQG	EKIIQEFLSK	VKQMPFTEMS	900
20	EENITIKLKQ	LKAEVIAKNN	SFVNEIISRI	KVTT			934

MSH2 (human cDNA) (SEQ ID NO:12)

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					ctctttggta	
					gttgatggcc	
					ctgtgtgaat	
					ggaccaaagg	
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					aaagacattt	
					agtgctgtat	
					aagtttttag	
					gacttcagcc	
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					acattaataa	
					tccagtgcac	
					aacaataaaa	
					aaattgactt	
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					ctcaatgatg	
					gcacctgttc	
					aaagcatcca	
					gtatactttg	
					aaatcaacat	
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10
    aaatgtcaga agaaaacatc acaataaagt taaaacagct aaaagctgaa gtaatagcaa 2820
    agaataatag ctttgtaaat gaaatcattt Cacgaataaa agttactacg tgaaaaatcc 2880
    cagtaatgga atgaaggtaa tattgataag ctattgtctg taatagtttt atattgtttt
     atattaaccc tttttccata gtgttaactg tcagtgccca tgggctatca acttaataag 3000
    atatttagta atattttact ttgaggacat tttcaaagat ttttattttg aaaaatgaga 3060
15
    gotgtaactg aggactgttt goaattgaca taggcaataa taagtgatgt gotgaatttt 3120
     ataaataaaa tcatgtagtt tgtgg
```

MLH1 (human) (SEO ID NO:13)

20	IQDNGTGIRK	EDLDIVCERF	TTSKLQSFED	LASISTYGFR	GEALASISHV	VKEGGLKLIQ AHVTITTKTA	120
						EEYGKILEVV IGCEDKTLAF	
	KMNGYISNAN	YSVKKCIFLL	FINHRLVEST	SLRKAIETVY	AAYLPKNTHP	FLYLSLEISP	300
25						GLAGPSGEMV IVTEDKTDIS	
23						KRHREDSDVE	
						VGCVNPQWAL	
						DSPESGWTEE EGLPIFILRL	
30						SWKWTVEHIV	
	YKALRSHILP	PKHFTEDGNI	LQLANLPDLY	KVFERC			756

MLH1 (human) (SEO ID NO:14)

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cttggctctt ctggcgccaa aatgtcgttc gtggcagggg ttattcggcg gctggacgag 60
acagtggtga acegeategc ggegggggaa gttatccagc ggccagctaa tgctatcaaa 120
gagatgattg agaactgttt agatgcaaaa tocacaagta ttcaagtgat tgttaaaqaq 180
qqaqqcctqa aqttqattca qatccaaqac aatqqcaccq qqatcaqqaa aqaaqatctq 240
gatattgtat gigaaaggtt cactactagt aaacigcagi cettigagga tittagecagi 300
atttetacet atggettteg aggtgagget ttggecagea taagecatgt ggetcatgtt 360
actattacaa cqaaaacaqc tqatqqaaaq tqtqcataca gagcaagtta ctcagatgga 420
aaactgaaag ceettetaa accatgtget ggcaatcaag ggacccagat cacggtggag 480
gacctttttt acaacatage cacgaggaga aaagetttaa aaaatecaag tgaagaatat 540
gggaaaattt tggaagttgt tggcaggtat tcagtacaca atgcaggcat tagtitctca 600
gttaaaaaac aaggagagac agtagctgat gttaggacac tacccaatgc ctcaaccgtg 660
gacaatatte getecatett tggaaatget gttagtegag aactgataga aattggatgt 720
gaggataaaa cectageett caaaatgaat ggttacatat ccaatgcaaa ctactcagtg 780
aagaagtgca tettettaet etteateaac eategtetgg tagaateaac tteettgaga 840
aaagccataq aaacagtgta tgcagcctat ttgcccaaaa acacacaccc attcctgtac 900
ctcaqtttaq aaatcaqtcc ccagaatgtg gatgttaatg tgcaccccac aaagcatgaa 960
gttcacttcc tgcacgagga gagcatcctg gagcgggtgc agcagcacat cgagagcaag 1020
ctcctgggct ccaattcctc caggatgtac ttcacccaga ctttgctacc aggacttgct 1080
ggcecctctg gggagatggt taaatccaca acaagtctga cotcgtcttc tacttctgga 1140
agtagtgata aggtetatge ccaccagatg gttcgtacag attcccggga acagaagett 1200
gatgeattte tgeagectet gageaaaece etgteeagte ageeceagge cattgteaca 1260
gaggataaga cagatatttc tagtggcagg gctaggcagc aagatgagga gatgcttgaa 1320
cteccaqcee etqetqaaqt ggetgecaaa aatcagaget tggaggggga tacaacaaaq 1380
gggacttcag aaatgtcaga gaagagaga cetacttcca gcaaccccag aaagagacat 1440
cqqqaaqatt ctqatqtqqa aatqqtqqaa gatqattccc qaaaqgaaat gactgcagct 1500
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5	aatgagcagg gtgaatcete aagettagtg ctcaggttat gagagtgget tttctgaaga	gacatgaggt agtgggcett aagaactgtt eggagecage ggacagagga agaaggetga	tctccgggag ggcacagcat ctaccagata accgctcttt agatggtccc gatgcttgca	atgttgcata caaaccaagt ctcatttatg gaccttgcca aaagaaggac gactatttct	accactcott tatacettet attttgccaa tgettgcett ttgetgaata etttggaaat	ggaagaaatt cgtgggctgt caacaccacc ttttggtgtt agatagtcca cattgttgag tgatgaggaa	1620 1680 1740 1800 1860 1920
10	cctatcttca gaaagcctca gagtcgaccc tggactgtgg	ttcttcgact gtaaagaatg tctcaggcca aacacattgt	agccactgag cgctatgttc gcagagtgaa ctataaagcc	gtgaattggg tattccatcc gtgcctggct ttgcgctcac	acgaagaaaa ggaagcagta ccattccaaa acattctgcc	ggagggactg ggaatgtttt catatctgag ctcctggaag tcctaaacat	2040 2100 2160 2220
15	gagaggtgtt cgatacaaag cacttaagac	aaatatggtt tgttgtatca	atttatgcac aagtgtgata cttctgatag	tgtgggatgt tacaaagtgt	gttcttcttt accaacataa	caaagtettt etetgtatte gtgttggtag attgattata	2340 2400

hPMS2-134 (human) (SEQ ID NO:15)

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20 MERABSSTE PAKAIKPIDR KSVHOICSGO VVLSLSTAVK ELVENSLDAG ATNIDLKLKD 60 YGYDLIEVSD NGCGVEEENF EGLTLKHHTS KIQEFADLTQ VETFGFRGEA LSSLCALSDV 120 TISTCHASAK VGT 133
```

hPMS2-134 (human cDNA) (SEQ ID NO:16)

25 cgaggcggat cgggtgttgc atccatggag cgagctgaga gctcgagtac agaacctgct 60 aaggccatca aacctattga tcggaagtca gtcactcaga tttgctctgg caggtggta 120 ctgagtctaa gcactgcggt aaaggagtta gtagaaaaca gtctgagtgc tggtgcact 180 aatattgatc taaagcttaa ggactatgga gtggatctta ttgaagttc agacaatgga 240 tgtggggtag aagaaagaaac ttctgaaggc ttaacctctga aacatcacaa atctaagatt 300 caagagtttg ccgacctaac tcaggtgaa actttaggct ttcgggggga agctctgagc 360 tcactttgtg cactgagcg tgtcaccatt tctacctgcc acgcatcggc gaaggttgg 420 acttga

For further information on the background of the invention the following references may be consulted, each of which is incorporated herein by reference in its entirety:

- Glaser, V. (1996) Can ReoPro repolish tarnished monoclonal therapeutics? Nat. Biotechol. 14:1216-1217.
- Weiner, L.M. (1999) Monoclonal antibody therapy of cancer. Semin. Oncol. 26:43 51.
- Saez-Llorens, X.E. et al. (1998) Safety and pharmacokinetics of an intramuscular humanized monoclonal antibody to respiratory syncytial virus in premature infants and infants with bronchopulmonary dysplasia. Pediat. Infect. Dis. J. 17:787-791.
- Shield, C.F. et al. (1996) A cost-effective analysis of OKT3 induction therapy in
 cadaveric kidney transplantation. Am. J. Kidney Dis. 27:855-864.

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- Khazaeli, M.B. et al. (1994) Human immune response to monoclonal antibodies. J. Immunother. 15:42-52.
- Emery, S.C. and W.J. Harris "Strategies for humanizing antibodies" In: ANTIBODY ENGINEERING C.A.K. Borrebaeck (Ed.) Oxford University Press, N.Y. 1995, pp. 159-183.
- U.S. Patent No. 5,530,101to Oueen and Selick.
- Reff, M.E. (1993) High-level production of recombinant immunoglobulins in mammalian cells. Curr. Opin. Biotechnol. 4:573-576.
- Neuberger, M. and M. Gruggermann, (1997) Monoclonal antibodies. Mice perform a
 human repertoire. Nature 386:25-26.
 - Fiedler, U. and U. Conrad (1995) High-level production and long-term storage of engineered antibodies in transgenic tobacco seeds. *Bio/Technology* 13:1090-1093.
 - Baker S.M. et al. (1995) Male defective in the DNA mismatch repair gene PMS2 exhibit abnormal chromosome synapsis in meiosis. Cell 82:309-319.
- 15 12. Bronner, C.E. et al. (1994) Mutation in the DNA mismatch repair gene homologue hMLHI is associated with hereditary non-polyposis colon cancer. Nature 368:258-261.
 - de Wind N. et al. (1995) Inactivation of the mouse Msh2 gene results in mismatch repair deficiency, methylation tolerance, hyperrecombination, and predisposition to cancer. Cell 82:321-300.
 - Drummond, J.T. et al. (1995) Isolation of an hMSH2-p160 heterodimer that restores mismatch repair to tumor cells. Science 268:1909-1912.
 - Modrich, P. (1994) Mismatch repair, genetic stability, and cancer. Science 266:1959-1960.
- Nicolaides, N.C. et al. (1998) A Naturally Occurring hPMS2 Mutation Can Confer a Dominant Negative Mutator Phenotype. Mol. Cell. Biol. 18:1635-1641.
 - Prolla, T.A. et al. (1994) MLH1, PMS1, and MSH2 Interaction during the initiation of DNA mismatch repair in yeast. Science 264:1091-1093.
- Strand, M. et al. (1993) Destabilization of tracts of simple repetitive DNA in yeast by
 mutations affecting DNA mismatch repair. Nature 365:274-276.

- Su, S.S., R.S. Lahue, K.G. Au, and P. Modrich (1988) Mispair specificity of methyl directed DNA mismatch corrections in vitro. J. Biol. Chem. 263:6829-6835.
- Parsons, R. et al. (1993) Hypermutability and mismatch repair deficiency in RER* tumor cells. Cell 75:1227-1236.
- Papadopoulos, N. et al. (1993) Mutation of a mutL homolog is associated with hereditary colon cancer. Science 263:1625-1629.
 - Perucho, M. (1996) Cancer of the microsatellite mutator phenotype. *Biol. Chem.* 377:675-684.
- Nicolaides N.C., K.W. Kinzler, and B. Vogelstein (1995) Analysis of the 5' region of
 PMS2 reveals heterogenous transcripts and a novel overlapping gene. Genomics
 29:329-334.
 - Nicolaides, N.C. et al. (1995) Genomic organization of the human PMS2 gene family. Genomics 30:195-206.
 - 25. Palombo, F. et al. (1994) Mismatch repair and cancer. Nature 36:417.
- Eshleman J.R. and S.D. Markowitz (1996) Mismatch repair defects in human carcinogenesis. *Hum. Mol. Genet.* 5:1489-494.
 - Liu, T. et al. (2000) Microsatellite instability as a predictor of a mutation in a DNA mismatch repair gene in familial colorectal cancer. Genes Chromosomes Cancer 27:17-25.
- Nicolaides, N.C. et al. (1992) The Jun family members, c-JUN and JUND, transactivate the human c-myb promoter via an Ap1 like element. J. Biol. Chem. 267:19665-19672.
 - Shields, R.L. et al. (1995) Anti-IgE monoclonal antibodies that inhibit allergenspecific histamine release. Int. Arch. Allergy Immunol. 107:412-413.
- Frigerio L. et al. (2000) Assembly, secretion, and vacuolar delivery of a hybrid immunoglobulin in plants. Plant Physiol. 123:1483-1494.
 - Bignami M, (2000) Unmasking a killer: DNA O(6)-methylguanine and the cytotoxicity of methylating agents. Mutat. Res. 462:71-82.

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- Drummond, J.T. et al. (1996) Cisplatin and adriamycin resistance are associated with MutLa and mismatch repair deficiency in an ovarian tumor cell line. J. Biol. Chem. 271:9645-19648.
- Galio, L. et al. (1999) ATP hydrolysis-dependent formation of a dynamic ternary nucleoprotein complex with MutS and MutL. Nucl. Acids Res. 27:2325-23231.

The above disclosure generally describes the present invention. A more complete understanding can be obtained by reference to the following specific examples which are provided herein for purposes of illustration only, and are not intended to limit the scope of the invention.

EXAMPLE 1: Stable expression of dominant negative MMR genes in hybridoma cells

It has been previously shown by Nicolaides et al. (Nicolaides et al. (1998) A Naturally Occurring hPMS2 Mutation Can Confer a Dominant Negative Mutator Phenotype Mol. Cell. Biol. 18:1635-1641) that the expression of a dominant negative allele in an otherwise MMR proficient cell could render these host cells MMR deficient. The creation of MMR deficient cells can lead to the generation of genetic alterations throughout the entire genome of a host organisms offspring, yielding a population of genetically altered offspring or siblings that may produce biochemicals with altered properties. This patent application teaches of the use of dominant negative MMR genes in antibody-producing cells, including but not limited to rodent hybridomas, human hybridomas, chimeric rodent cells producing human immunoglobulin gene products, human cells expressing immunoglobulin genes, mammalian cells producing single chain antibodies, and prokaryotic cells producing mammalian immunoglobulin genes or chimeric immunoglobulin molecules such as those contained within single-chain antibodies. The cell expression systems described above that are used to produce antibodies are well known by those skilled in the art of antibody therapeutics.

To demonstrate the ability to create MMR defective hybridomas using dominant negative alleles of MMR genes, we first transfected a mouse hybridoma cell line that is known to produce and antibody directed against the human IgE protein with an expression vector containing the human PMS2 (cell line referred to as HBPMS2), the previously

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published dominant negative PMS2 mutant referred herein as PMS134 (cell line referred to as HB134), or with no insert (cell line referred to as HBvec). The results showed that the PMS134 mutant could indeed exert a robust dominant negative effect, resulting in biochemical and genetic manifestations of MMR deficiency. Unexpectedly was the finding that the full length PMS2 also resulted in a lower MMR activity while no effect was seen in cells containing the empty vector. A brief description of the methods is provided below.

The MMR proficient mouse H36 hybridoma cell line was transfected with various hPMS2 expression plasmids plus reporter constructs for assessing MMR activity. The MMR genes were cloned into the pEF expression vector, which contains the elongation factor promoter upstream of the cloning site followed by a mammalian polyadenvlation signal. This vector also contains the NEOr gene that allows for selection of cells retaining this plasmid. Briefly, cells were transfected with 1 µg of each vector using polyliposomes following the manufacturer's protocol (Life Technologies). Cells were then selected in 0.5 mg/ml of G418 for 10 days and G418 resistant cells were pooled together to analyze for gene expression. The pEF construct contains an intron that separates the exon 1 of the EF gene from exon 2, which is juxtaposed to the 5' end of the polylinker cloning site. This allows for a rapid reverse transcriptase polymerase chain reaction (RT-PCR) screen for cells expressing the spliced products. At day 17, 100,000 cells were isolated and their RNA extracted using the trizol method as previously described (Nicolaides N.C., Kinzler, K.W., and Vogelstein, B. (1995) Analysis of the 5' region of PMS2 reveals heterogeneous transcripts and a novel overlapping gene. Genomics 29:329-334). RNAs were reverse transcribed using Superscript II (Life Technologies) and PCR amplified using a sense primer located in exon 1 of the EF gene (5'-ttt cgc aac ggg ttt gcc g-3') and an antisense primer (5'-gtt tca gag tta agc ctt cg-3') centered at nt 283 of the published human PMS2 cDNA, which will detect both the full length as well as the PMS134 gene expression. Reactions were carried out using buffers and conditions as previously described (Nicolaides, N.C., et al. (1995) Genomic organization of the human PMS2 gene family. Genomics 30:195-206), using the following amplification parameters: 94°C for 30 sec, 52°C for 2 min, 72°C for 2 min, for 30 cycles. Reactions were analyzed on agarose gels. Figure 1 shows a representative example of PMS expression in stably transduced H36 cells.

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Expression of the protein encoded by these genes were confirmed via western blot using a polyclonal antibody directed to the first 20 amino acids located in the N-terminus of the protein following the procedures previously described (data not shown) (Nicolaides et al. (1998) A Naturally Occurring hPMS2 Mutation Can Confer a Dominant Negative Mutator Phenotype. Mol. Cell. Biol. 18:1635-1641.

EXAMPLE 2: hPMS134 Causes a Defect in MMR Activity and hypermutability in hybridoma cells

A hallmark of MMR deficiency is the generation of unstable microsatellite repeats in the genome of host cells. This phenotype is referred to as microsatellite instability (MI) (Modrich, P. (1994) Mismatch repair, genetic stability, and cancer Science 266:1959-1960; Palombo, F., et al. (1994) Mismatch repair and cancer Nature 36:417). MI consists of deletions and/or insertions within repetitive mono-, di- and/or tri nucleotide repetitive sequences throughout the entire genome of a host cell. Extensive genetic analysis eukaryotic cells have found that the only biochemical defect that is capable of producing MI is defective MMR (Strand, M., et al. (1993) Destabilization of tracts of simple repetitive DNA in yeast by mutations affecting DNA mismatch repair Nature 365:274-276; Perucho, M. (1996) Cancer of the microsatellite mutator phenotype. Biol Chem. 377:675-684; Eshleman J.R., and Markowitz, S.D. (1996) Mismatch repair defects in human carcinogenesis. Hum. Mol. Genet. 5:1489-494). In light of this unique feature that defective MMR has on promoting MI, it is now used as a biochemical marker to survey for lack of MMR activity within host cells (Perucho, M. (1996) Cancer of the microsatellite mutator phenotype. Biol Chem. 377:675-684; Eshleman J.R., and Markowitz, S.D. (1996) Mismatch repair defects in human carcinogenesis. Hum. Mol. Genet. 5:1489-494; Liu, T., et al. (2000) Microsatellite instability as a predictor of a mutation in a DNA mismatch repair gene in familial colorectal cancer Genes Chromosomes Cancer 27:17-25).

A method used to detect MMR deficiency in eukaryotic cells is to employ a reporter gene that has a polynucleotide repeat inserted within the coding region that disrupts its reading frame due to a frame shift. In the case where MMR is defective, the reporter gene will acquire random mutations (i.e. insertions and/or deletions) within the polynucleotide

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repeat yielding clones that contain a reporter with an open reading frame. We have employed the use of an MMR-sensitive reporter gene to measure for MMR activity in HByec. HBPMS2, and HBPMS134 cells. The reporter construct used the pCAR-OF, which contains a hygromycin resistance (HYG) gene plus a β-galactosidase gene containing a 29 bp out-of-frame poly-CA tract at the 5' end of its coding region. The pCAR-OF reporter would not generate β -galactosidase activity unless a frame-restoring mutation (i.e., insertion or deletion) arose following transfection. HBvec, HBPMS2, and HB134 cells were each transfected with pCAR-OF vector in duplicate reactions following the protocol described in Example 1. Cells were selected in 0.5 mg/ml G418 and 0.5mg/ml HYG to select for cells retaining both the MMR effector and the pCAR-OF reporter plasmids. All cultures transfected with the pCAR vector resulted in a similar number of HYG/G418 resistant cells. Cultures were then expanded and tested for \beta-galactosidase activity in situ as well as by biochemical analysis of cell extracts. For in situ analysis, 100,000 cells were harvested and fixed in 1% gluteraldehyde, washed in phosphate buffered saline solution and incubated in 1 ml of X-gal substrate solution [0.15 M NaCl, 1 mM MgCl₂, 3.3 mM K₄Fe(CN)₆, 3.3 mM K₃Fe(CN)₆, 0.2% X-Gal] in 24 well plates for 2 hours at 37°C. Reactions were stopped in 500 mM sodium bicarbonate solution and transferred to microscope slides for analysis. Three fields of 200 cells each were counted for blue (β-galactosidase positive cells) or white (β-galactosidase negative cells) to assess for MMR inactivation. Table 1 shows the results from these studies. While no β-galactosidase positive cells were observed in HBvec cells, 10% of the cells per field were β -galactosidase positive in HB134 cultures and 2% of the cells per field were β-galactosidase positive in HBPMS2 cultures.

Cell extracts were prepared from the above cultures to measure β-galactosidase using a quantitative biochemical assay as previously described (Nicolaides *et al.* (1998)

25 A Naturally Occurring hPMS2 Mutation Can Confer a Dominant Negative Mutator Phenotype *Mol. Cell. Biol.* 18:1635-1641; Nicolaides, N.C., *et al.* (1992) The Jun family members, c-JUN and JUND, transactivate the human *c-myb* promoter via an Ap1 like element. *J. Biol. Chem.* 267:19665-19672). Briefly, 100,000 cells were collected, centrifuged and resuspended in 200 μls of 0.25M Tris, pH 8.0. Cells were

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lysed by freeze/thawing three times and supernatants collected after microfugation at 14,000 rpms to remove cell debris. Protein content was determined by spectrophotometric analysis at OD^{280} . For biochemical assays, $20~\mu g$ of protein was added to buffer containing 45 mM 2-mercaptoethanol, $1 mM \, MgCl_2$, $0.1~M \, NaPO_4$ and 0.6~mg/ml Chlorophenol red- β -D-galactopyranoside (CPRG, Boehringer Mannheim). Reactions were incubated for 1 hour, terminated by the addition of $0.5~M \, Na_2CO_3$, and analyzed by spectrophotometry at 576 nm. H36 cell lysates were used to subtract out background. Figure 2 shows the β -galactosidase activity in extracts from the various cell lines. As shown, the HB134 cells produced the highest amount of β -galactosidase, while no activity was found in the HBvec cells containing the pCAR-OF. These data demonstrate the ability to generate MMR defective hybridoma cells using dominant negative MMR gene alleles.

 Table 1. β-galactosidase expression of HBvec, HBPMS2 and HB134 cells transfected with pCAR-OF reporter vectors. Cells were transfected with the pCAR-OF β-galactosidase reporter plasmid. Transfected cells were selected in hygromycin and G418, expanded and stained with X-gal solution to measure for β -galactosidase activity (blue colored cells). 3 fields of 200 cells each were analyzed by microscopy. The results below represent the mean +/- standard deviation of these experiments.

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Table 1.

CELL LINE	# BLUE CELLS
HBvec	0 +/- 0
HBPMS2	4 +/- 1
HB134	20 +/- 3

EXAMPLE 3: Screening strategy to identify hybridoma clones producing antibodies with higher binding affinities and/or increased immunoglobulin production.

An application of the methods presented within this document is the use of MMR. deficient hybridomas or other immunoglobulin producing cells to create genetic alterations within an immunoglobulin gene that will yield antibodies with altered biochemical properties. An illustration of this application is demonstrated within this example whereby the HB134 hybridoma (see Example 1), which is a MMR-defective cell line that produces an anti-human immunoglobulin type E (hIgE) MAb, is grown for 20 generations and clones are isolated in 96-well plates and screened for hIgE binding. Figure 3 outlines the screening procedure to identify clones that produce high affinity MAbs, which is presumed to be due to an alteration within the light or heavy chain variable region of the protein. The assay employs the use of a plate Enzyme Linked Immunosorbant Assay (ELISA) to screen for clones that produce highaffinity MAbs. 96-well plates containing single cells from HBvec or HB134 pools are grown for 9 days in growth medium (RPMI 1640 plus 10% fetal bovine serum) plus 0.5 mg/ml G418 to ensure clones retain the expression vector. After 9 days, plates are screened using an hIgE plate ELISA, whereby a 96 well plate is coated with 50µls of a 1µg/ml hIgE solution for 4 hours at 4°C. Plates are washed 3 times in calcium and magnesium free phosphate buffered saline solution (PBS-1) and blocked in 100uls of PBS-1 with 5% dry milk for 1 hour at room temperature. Wells are rinsed and incubated with 100 µls of a PBS solution containing a 1:5 dilution of conditioned medium from each cell clone for 2 hours. Plates are then washed 3 times with PBS+ and incubated for 1 hour at room temperature with 50 µls of a PBS+ solution containing 1:3000 dilution of a sheep anti-mouse horse radish peroxidase (HRP) conjugated

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secondary antibody. Plates are then washed 3 times with PBS+ and incubated with 50 µls of TMB-HRP substrate (BioRad) for 15 minutes at room temperature to detect amount of antibody produced by each clone. Reactions are stopped by adding 50 µls of 500mM sodium bicarbonate and analyzed by OD at 415nm using a BioRad plate reader. Clones exhibiting an enhanced signal over background cells (H36 control cells) are then isolated and expanded into 10 ml cultures for additional characterization and confirmation of ELISA data in triplicate experiments. ELISAs are also performed on conditioned (CM) from the same clones to measure total Ig production within the conditioned medium of each well. Clones that produce an increased ELISA signal and have increased antibody levels are then further analyzed for variants that over-express and/or over-secrete antibodies as described in Example 4. Analysis of five 96-well plates each from HBvec or HB134 cells have found that a significant number of clones with a higher Optical Density (OD) value is observed in the MMR-defective HB134 cells as compared to the HByec controls. Figure 4 shows a representative example of HB134 clones producing antibodies that bind to specific antigen (in this case IgE) with a higher affinity. Figure 4 provides raw data from the analysis of 96 wells of HBvec (left graph) or HB134 (right graph) which shows 2 clones from the HB134 plate to have a higher OD reading due to 1) genetic alteration of the antibody variable domain that leads to an increased binding to IgE antigen, or 2) genetic alteration of a cell host that leads to over-production/secretion of the antibody molecule. Anti-Ig ELISA found thatthe two clones, shown in figure 4 have Ig levels within their CM similar to the surrounding wells exhibiting ower OD values. These data suggest that a genetic alteration occurred within the antigen binding domain of the antibody which in turn allows for higher binding to antigen.

Clones that produced higher OD values as determined by ELISA were further analyzed at the genetic level to confirm that mutations within the light or heavy chain variable region have occurred that lead to a higher binding affinity hence yielding to a stronger ELISA signal. Briefly, 100,000 cells are harvested and extracted for RNA using the Triazol method as described above. RNAs are reverse transcribed using Superscript II as suggested by the manufacturer (Life Technology) and PCR amplified for the antigen binding sites contained within the variable light and heavy chains. Because of the heterogeneous nature of these

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genes, the following degenerate primers are used to amplify light and heavy chain alleles from the parent H36 strain.

Light chain sense: 5'-GGA TTT TCA GGT GCA GAT TTT CAG-3' (SEQ ID NO:1)

Light chain antisense: 5'-ACT GGA TGG TGG GAA GAT GGA-3' (SEQ ID NO:2)

Heavy chain sense: 5'-A(G/T) GTN (A/C)AG CTN CAG (C/G)AG TC-3' (SEQ ID NO:3)

Heavy chain antisense: 5'-TNC CTT G(A/G)C CCC AGT A(G/A)(A/T)C-3' (SEQ ID NO:4)

PCR reactions using degenerate oligonucleotides are carried out at 94°C for 30 sec, 52°C for 1 min, and 72°C for 1 min for 35 cycles. Products are analyzed on agarose gels. Products of the expected molecular weights are purified from the gels by Gene Clean (Bio 101), cloned into T-tailed vectors, and sequenced to identify the wild type sequence of the variable light and heavy chains. Once the wild type sequence has been determined, nondegenerate primers were made for RT-PCR amplification of positive HB134 clones. Both the light and heavy chains were amplified, gel purified and sequenced using the corresponding sense and antisense primers. The sequencing of RT-PCR products gives representative sequence data of the endogenous immunoglobulin gene and not due to PCR induced mutations. Sequences from clones were then compared to the wild type sequence for sequence comparison. An example of the ability to create in vivo mutations within an immunoglobulin light or heavy chain is shown in figure 5, where HB134 clone92 was identified by ELISA to have an increased signal for hIgE. The light chain was amplified using specific sense and antisense primers. The light chain was RT-PCR amplified and the resulting product was purified and analyzed on an automated ABI377 sequencer. As shown in clone A, a residue -4 upstream of the CDR region 3 had a genetic change from ACT to TCT, which results in a Thr to Ser change within the framework region just preceding the CDR#3. In clone B, a residue -6 upstream of the CDR region had a genetic change from CCC to CTC. which reslts in a Pro to His change within framework region preceeding CDR#2.

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The ability to generate random mutations in immunoglobulin genes or chimeric immunoglobulin genes is not limited to hybridomas. Nicolaides et al. (Nicolaides et al. (1998) A Naturally Occurring hPMS2 Mutation Can Confer a Dominant Negative Mutator Phenotype Mol. Cell. Biol. 18:1635-1641) has previously shown the ability to generate hypermutable hamster cells and produce mutations within an endogenous gene. A common method for producing humanized antibodies is to graft CDR sequences from a MAb (produced by immunizing a rodent host) onto a human Ig backbone, and transfection of the chimeric genes into Chinese Hamster Ovary (CHO) cells whih in turn produce a functional Ab that is secreted by the CHO cells (Shields, R.L., et al. (1995) Anti-IgE monoclonal antibodies that inhibit allergen-specific histamine release. Int. Arch. Allergy Immunol. 107:412-413). The methods described within this application are also useful for generating genetic alterations within Ig genes or chimeric Igs transfected within host cells such as rodent cell lines, plants, yeast and prokaryotes (Frigerio L, et al. (2000) Assembly, secretion, and vacuolar delivery of a hybrid immunoglobulin in plants. Plant Physiol. 123:1483-1494).

These data demonstrate the ability to generate hypermutable hybridomas, or other Ig producing host cells that can be grown and selected, to identify structurally altered immunoglobulins yielding antibodies with enhanced biochemical properties, including but not limited to increased antigen binding affinity. Moreover, hypermutable clones that contain missense mutations within the immunoglobulin gene that result in an amino acid change or changes can be then further characterized for *in vivo* stability, antigen clearance, on-off binding to antigens, etc. Clones can also be further expanded for subsequent rounds of *in vivo* mutations and can be screened using the strategy listed above.

The use of chemical mutagens to produce genetic mutations in cells or whole organisms are limited due to the toxic effects that these agents have on "normal" cells. The use of chemical mutagens such as MNU in MMR defective organisms is much more tolerable yielding to a 10 to 100 fold increase in genetic mutation over MMR deficiency alone (Bignami M, (2000) Unmasking a killer: DNA O(6)-methylguanine and the cytotoxicity of methylating agents. *Mutat. Res.* 462:71-82). This strategy allows for the use of chemical mutagens to be used in MMR-defective Ab producing cells as a method for increasing

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additional mutations within immunoglobulin genes or chimeras that may yield functional Abs with altered biochemical properties such as enhanced binding affinity to antigen, etc.

Example 4: Generation of antibody producing cells with enhanced antibody production

Analysis of clones from H36 and HB134 following the screening strategy listed above hasidentified a significant number of clones that produce enhanced amounts of antibody into the medium. While a subset of these clones gave higher Ig binding data as determined by ELISA as a consequence of mutations within the antigen binding domains contained in the variable regions, others were found to contain "enhanced" antibody production. A summary of the clones producing enhanced amounts of secreted MAb is shown in TABLE 2, where a significant number of clones from HB134 cells were found to produce enhanced Ab production within the conditioned medium as compared to H36 control cells.

TABLE 2. Generation of hybridoma cells producing high levels of antibody. HB134 clones were assayed by ELISA for elevated Ig levels. Analysis of 480 clones showed that a significant number of clones had elevated MAb product levels in their CM. Quantification showed that several of these clones produced greater than 500ngs/ml of MAb due to either enhanced expression and/or secretion as compared to clones from the H36 cell line.

20 Table 2. Production of MAb in CM from H36 and HB134 clones.

Cell Line	% clones > 400 ng/ml	% clones >500 ng/ml
H36	1/480 = 0.2%	0/480 = 0%
HB134	50/480 = 10%	8/480 = 1.7%

Cellular analysis of HB134 clones with higher MAb levels within the conditioned medium (CM) were analyzed to determine if the increased production was simply due to genetic alterations at the Ig locus that may lead to over-expression of the polypeptides forming the antibody, or due to enhanced secretion due to a genetic alteration affecting secretory pathway mechanisms. To address this issue, we expanded three HB134 clones that had increased levels of antibody within their CM. 10,000 cells were prepared for western blot

elevated secretion of antibody.

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analysis to assay for intracellular steady state Ig protein levels (Figure 6). In addition, H36 cells were used as a standard reference (Lane 2) and a rodent fibroblast (Lane 1) was used as an Ig negative control. Briefly, cells were pelleted by centrifugation and lysed directly in 300 μl of SDS lysis buffer (60 mM Tris, pH 6.8, 2% SDS, 10% glycerol, 0.1 M 2-mercaptoethanol, 0.001% bromophenol blue) and boiled for 5 minutes. Lysate proteins were separated by electrophoresis on 4-12% NuPAGE gels (for analysis of Ig heavy chain, Gels were electroblotted onto Immobilon-P (Millipore) in 48 mM Tris base, 40 mM glycine, 0.0375% SDS, 20% methanol and blocked at room temperature for 1 hour in Tris-buffered saline (TBS) plus 0.05% Tween-20 and 5% condensed milk. Filters were probed with a 1:10,000 dilution of sheep anti-mouse horseradish peroxidase conjugated monoclonal antibody in TBS buffer and detected by chemiluminescence using Supersignal substrate (Pierce). Experiments were repeated in duplicates to ensure reproducibility. Figure 6 shows a representative analysis where a subset of clones had enhanced Ig production which accounted for increased Ab production (Lane 5) while others had a similar steady state level as the control sample, yet had higher levels of Ab within the CM. These data suggest a mechanism whereby a subset of HB134 clones contained a genetic alteration that in turn produces

The use of chemical mutagens to produce genetic mutations in cells or whole organisms are limited due to the toxic effects that these agents have on "normal" cells. The use of chemical mutagens such as MNU in MMR defective organisms is much more tolerable yielding to a 10 to 100 fold increase in genetic mutation over MMR deficiency alone (Bignami M, (2000) Unmasking a killer: DNA O(6)-methylguanine and the cytotoxicity of methylating agents. *Mutat. Res.* 462:71-82). This strategy allows for the use of chemical mutagens to be used in MMR-defective Ab producing cells as a method for increasing additional mutations within immunoglobulin genes or chimeras that may yield functional Abs with altered biochemical properties such as enhanced binding affinity to antigen, etc.

Example 5: establishment of genetic stability in hybridoma cells with new output trait.

The initial steps of MMR are dependent on two protein complexes, called MutSα and MutLα (Nicolaides *et al.* (1998) A Naturally Occurring hPMS2 Mutation Can Confer a

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Dominant Negative Mutator Phenotype. *Mol. Cell. Biol.* 18:1635-1641). Dominant negative MMR alleles are able to perturb the formation of these complexes with downstream biochemicals involved in the excision and polymerization of nucleotides comprising the "corrected" nucleotides. Examples from this application show the ability of a truncated MMR allele (PMS134) as well as a full length human PMS2 when expressed in a hybridoma cell line is capable of blocking MMR resulting in a hypermutable cell line that gains genetic alterations throughout its entire genome per cell division. Once a cell line is produced that contains genetic alterations within genes encoding for an antibody, a single chain antibody, over expression of immunoglobulin genes and/or enhanced secretion of antibody, it is desirable to restore the genomic integrity of the cell host. This can be achieved by the use of inducible vectors whereby dominant negative MMR genes are cloned into such vectors, introduced into Ab producing cells and the cells are cultured in the presence of inducer molecules and/or conditions. Inducible vectors include but are not limited to chemical regulated promoters such as the steroid inducible MMTV, tetracycline regulated promoters, temperature sensitive MMR gene alleles, and temperature sensitive promoters.

The results described above lead to several conclusions. First, expression of hPMS2 and PMS134 results in an increase in microsatellite instability in hybridoma cells. That this elevated microsatellite instability is due to MMR deficiency was proven by evaluation of extracts from stably transduced cells. The expression of PMS134 results in a polar defect in MMR, which was only observed using heteroduplexes designed to test repair from the 5' direction (no significant defect in repair from the 3' direction was observed in the same extracts) (Nicolaides et al. (1998) A Naturally Occurring hPMS2 Mutation Can Confer a Dominant Negative Mutator Phenotype. Mol. Cell. Biol. 18:1635-1641). Interestingly, cells deficient in hMLH1 also have a polar defect in MMR, but in this case preferentially affecting repair from the 3' direction (Drummond, J.T, et al. (1996) Cisplatin and adriamycin resistance are associated with MutLa and mismatch repair deficiency in an ovarian tumor cell line. J. Biol. Chem. 271:9645-19648). It is known from previous studies in both prokaryotes and eukaryotes that the separate enzymatic components mediate repair from the two different directions. Our results, in combination with those of Drummond et al. (Shields, R.L., et al. (1995) Anti-IgE monoclonal antibodies that inhibit allergen-specific histamine release. Int.

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Arch Allergy Immunol. 107:412-413), strongly suggest a model in which 5' repair is primarily dependent on hPMS2 while 3' repair is primarily dependent on hMLH1. It is easy to envision how the dimeric complex between PMS2 and MLH1 might set up this directionality. The combined results also demonstrate that a defect in directional MMR is sufficient to produce a MMR defective phenotype and suggests that any MMR gene allele is useful to produce genetically altered hybridoma cells, or a cell line that is producing Ig gene products. Moreover, the use of such MMR alleles will be useful for generating genetically altered Ig polypeptides with altered biochemical properties as well as cell hosts that produce enhanced amounts of antibody molecules.

Another method that is taught in this application is that ANY method used to block MMR can be performed to generate hypermutability in an antibody-producing cell that can lead to genetically altered antibodies with enhanced biochemical features such as but not limited to increased antigen binding, enhanced pharmacokinetic profiles, etc. These processes can also to be used to generate antibody producer cells that have increased Ig expression as shown in Example 4, figure 6 and/or increased antibody secretion as shown in Table 2.

In addition, we demonstrate the utility of blocking MMR in antibody producing cells to increase genetic alterations within Ig genes that may lead to altered biochemical features such as, but not limited to, increased antigen binding affinities (Figure 5A and 5B). The blockade of MMR in such cells can be through the use of dominant negative MMR gene alleles from any species including bacteria, yeast, protozoa, insects, rodents, primates, mammalian cells, and man. Blockade of MMR can also be generated through the use of antisense RNA or deoxynucleotides directed to any of the genes involved in the MMR biochemical pathway. Blockade of MMR can be through the use of polypeptides that interfere with subunits of the MMR complex including but not limited to antibodies. Finally, the blockade of MMR may be through the use chemicals such as but not limited to nonhydrolyzable ATP analogs, which have been shown to block MMR (Galio, L, et al. (1999) ATP hydrolysis-dependent formation of a dynamic ternary nucleoprotein complex with MutS and MutL. Nucl. Acids Res. 27:2325-23231).

Docket No.: MOR-0003

WE CLAIM:

- A method for making a hypermutable, antibody producing cell, comprising introducing into a cell capable of producing antibodies a polynucleotide comprising a dominant negative allele of a mismatch repair gene.
- The method of claim 1 wherein said polynucleotide is introduced by transfection of a suspension of cells in vitro.
- 3. The method of claim 1 wherein said mismatch repair gene is PMS2.
- 4. The method of claim 1 wherein said mismatch repair gene is human PMS2.
- 5. The method of claim 1 wherein said mismatch repair gene is MLH1.
- 6. The method of claim 1 wherein said mismatch repair gene is PMS1.
- 7. The method of claim 1 wherein said mismatch repair gene is MSH2.
- 8. The method of claim 1 wherein said mismatch repair gene is MSH2.
- 9. The method of claim 4 wherein said allele comprises a truncation mutation.
- The method of claim 4 wherein said allele comprises a truncation mutation at codon
 134.
- The method of claim 10 wherein said truncation mutation is a thymidine at nucleotide 424 of wild-type PMS2.
- The method of claim 1 wherein said polynucleotide is introduced into a fertilized egg of an animal.
- 13. The method of claim 12 wherein the fertilized egg is subsequently implanted into a pseudo-pregnant female whereby the fertilized egg develops into a mature transgenic animal.
- 14. The method of claim 12 wherein said mismatch repair gene is PMS2.
- 15. The method of claim 12 wherein said mismatch repair gene is human PMS2.
- 16. The method of claim 12 wherein said mismatch repair gene is human MLH1.
- 17. The method of claim 12 wherein said mismatch repair gene is human PMS1.
- 18. The method of claim 11 wherein said mismatch repair gene is a human mutL homolog.
- 19. The method of claim 15 wherein said allele comprises a truncation mutation.
- The method of claim 15 wherein said allele comprises a truncation mutation at codon 134.

- The method of claim 19 wherein said truncation mutation is a thymidine at nucleotide 424 of wild-type PMS2.
- The method of claim 1 wherein said capability is due to the co-introduction of an immunoglobulin gene into said cell.
- A homogeneous culture of hypermutable, mammalian cells wherein said cells comprise a dominant negative allele of a mismatch repair gene.
- The culture of hypermutable, mammalian cells of claim 23 wherein the mismatch repair gene is PMS2.
- The culture of hypermutable, mammalian cells of claim 24 wherein the mismatch repair gene is human PMS2.
- The culture of hypermutable, mammalian cells of claim 23 wherein the mismatch repair gene is MLH1.
- The culture of hypermutable, mammalian cells of claim 23 wherein the mismatch repair gene is PMSI.
- The culture of hypermutable, mammalian cells of claim 23 wherein the mismatch repair gene is a human mutL homolog.
- The culture of hypermutable, mammalian cells of claim 23 wherein the cells express a
 protein consisting of the first 133 amino acids of hPMS2.
- A method for generating a mutation in a gene affecting antibody production in an antibody-producing cell comprising:

growing a said cell comprising said gene and a dominant negative allele of a mismatch repair gene: and

testing the cell to determine whether said gene of interest harbors a mutation.

- 31. The method of claim 30 wherein the step of testing comprises analyzing a nucleotide sequence of said gene.
- The method of claim 30 wherein the step of testing comprises analyzing mRNA transcribed from said gene.
- 33. The method of claim 30 wherein the step of testing comprises analyzing a protein encoded by the gene of interest.

- 34. The method of claim 30 wherein the step of testing comprises analyzing the phenotype of said gene.
- The method of claim 30 wherein the step of testing comprises analyzing the binding activity of an antibody.
- 36. A method wherein a mammalian cell is made MMR defective by the process of introducing a polynucleotide comprising an antisense oligonucleotide targeted against an allele of a mismatch repair gene into a mammalian cell, whereby the cell becomes hypermutable.
- The method of claim 36 wherein the step of testing comprises analyzing a nucleotide sequence of said gene.
- The method of claim 36 wherein the step of testing comprises analyzing mRNA transcribed from said gene.
- The method of claim 36 wherein the step of testing comprises analyzing a protein encoded by said gene.
- 40. The method of claim 36 wherein the step of testing comprises analyzing the phenotype of said gene.
- The method of claim 36 wherein the step of testing comprises analyzing the binding activity of an antibody.
- A method for generating a mutation in a gene affecting antibody production in an antibody-producing cell comprising:

growing said cell comprising said gene and a polynucleotide encoding a dominant negative allele of a mismatch repair gene; and

testing said cell to determine whether said cell harbors at least one mutation in said gene yielding to a new biochemical feature to the product of said gene, wherein said new biochemical feature is selected from the group consisting of over-expression of said product, enhanced secretion of said product, enhanced affinity of said product for antigen, and combinations thereof.

43. The method of claim 42 wherein the step of testing comprises analyzing the steady state expression of the immunoglobulin gene of said cell.

- 44. The method of claim 42 wherein the step of testing comprises analyzing steady state mRNA transcribed from the immunoglobulin gene of said cell.
- 45. The method of claim 42 wherein the step of testing comprises analyzing the amount of secreted protein encoded by the immunoglobulin gene of said cell.
- 46. The method of claim 36 wherein the cell is made by the process of introducing a polynucleotide comprising a dominant negative allele of a mismatch repair gene into a cell in the presence of DNA mutagens.
- The method of claim 46 wherein the step of testing comprises analyzing a nucleotide sequence of an immunoglobulin gene of said cell.
- 48. The method of claim 46 wherein the step of testing comprises analyzing mRNA transcribed from the immunoglobulin gene of said cell.
- The method of claim 46 wherein the step of testing comprises analyzing the immunoglobulin protein encoded by said gene.
- The method of claim 46 wherein the step of testing comprises analyzing the biochemical activity of the protein encoded by said gene.
- 51. A hypermutable transgenic mammalian cell made by the method of claim 42.
- 52. The transgenic mammalian cell of claim 51 wherein said cell is from primate.
- 53. The transgenic mammalian cell of claim 51 wherein said cell is from rodent.
- 54. The transgenic mammalian cell of claim 51 wherein said cell is from human.
- 55. The transgenic mammalian cell of claim 51 wherein said cell is eukaryotic.
- 56. The transgenic mammalian cell of claim 51 wherein said cell is prokaryotic
- 57. A method of reversibly altering the hypermutability of an antibody producing cell comprising introducing an inducible vector into a cell, wherein said inducible vector comprises a dominant negative allele of a mismatch repair gene operably linked to an inducible promoter, and inducing said cell to express said dominant negative mismatch repair gene.
- 58. The method of claim 57 wherein said mismatch repair gene is PMS2.
- 59. The method of claim 58 wherein said mismatch repair gene is human PMS2.
- 60. The method of claim 57 wherein said mismatch repair gene is MLH1.
- 61. The method of claim 57 wherein said mismatch repair gene is PMS1.

- 62. The method of claim 57 wherein said mismatch repair gene is a human mutL homolog.
- 63. The method of claim 57 wherein said cell expresses a protein consisting of the first 133 amino acids of hPMS2.
- 64. The method of claim 57 further comprising analyzing the immunoglobulin protein expressed by said antibody producing cell.
- The method of claim 64 further comprising ceasing induction of said cell, thereby restoring genetic stability of said cell.
- 66. A method of producing genetically altered antibodies comprising

transfecting a polynucleotide encoding an immunoglobulin protein into a cell, wherein said cell comprises a dominant negative mismatch repair gene;

growing said cell, thereby producing a hypermutated polynucleotide encoding a hypermutated immunoglobulin protein;

isolating said hypermutated polynucleotide; and

transfecting said hypermutated polynucleotide into a genetically stable cell, thereby producing a hypermutated antibody-producing, genetically stable cell.

- 67. The method of claim 66 wherein said mismatch repair gene is PMS2.
- 68. The method of claim 66 wherein said mismatch repair gene is human PMS2.
- 69. The method of claim 66 wherein said mismatch repair gene is MLH1.
- 70. The method of claim 66 wherein said mismatch repair gene is *PMS1*.
- 71. The method of claim 66 wherein said mismatch repair gene is a human mutL homolog.
- The method of claim 66 wherein said cell expresses a protein consisting of the first
 133 amino acids of hPMS2.

ABSTRACT

Dominant negative alleles of human mismatch repair genes can be used to generate hypermutable cells and organisms. By introducing these genes into cells and transgenic animals, new cell lines and animal varieties with novel and useful properties can be prepared more efficiently than by relying on the natural rate of mutation. These methods are useful for generating genetic diversity within immunoglobulin genes directed against an antigen of interest to produce altered antibodies with enhanced biochemical activity. Moreover, these methods are useful for generating antibody-producing cells with increased level of antibody production.

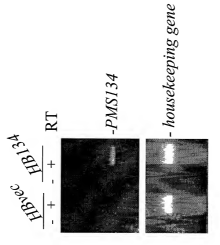


FIG. 1/6

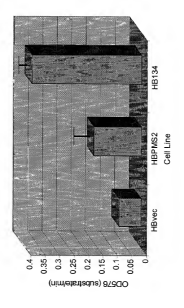


Fig. 2/(

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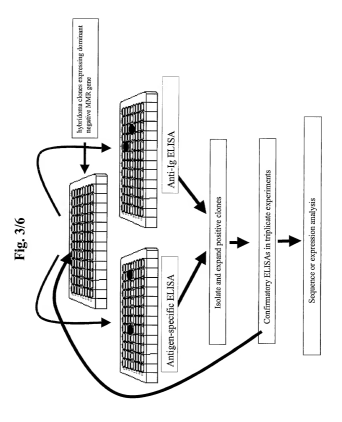
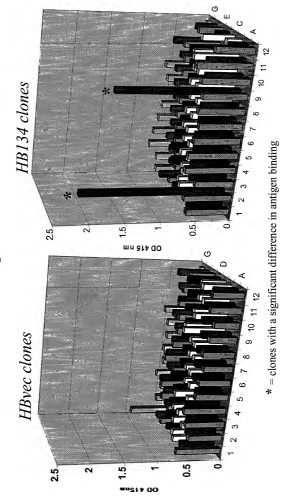


Fig.4/6



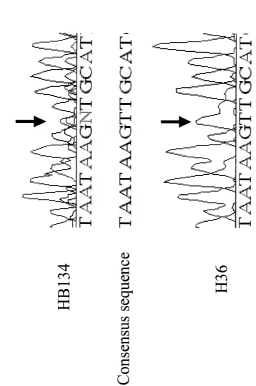
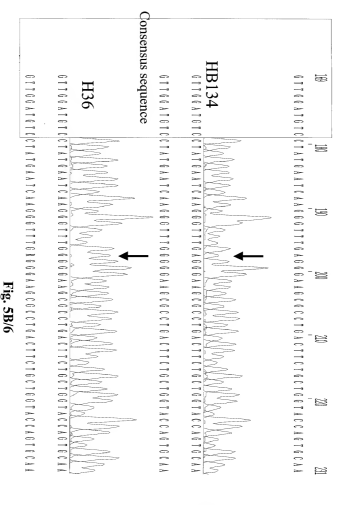


Fig. 5A/6

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Ig heavy chain

Fig 6/6

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In Re Application of

Nicholas C. Nicolaides, Luigi Grasso, and Philip M. Sass

Group Art Unit: Not assigned

Examiner: Not assigned

For: METHODS FOR GENERATING GENETICALLY ALTERED ANTIBODY-PRODUCING CELL LINES WITH IMPROVED ANTIBODY

CHARACTERISTICS

M

DECLARATION AND POWER OF ATTORNEY

As a below named inventor, I hereby declare that:

Utility Patent

My residence, post office address and citizenship are as stated below next to my name; and

I believe that I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a

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is sought on the inv	rention, whose title app	pears above, the specification of which:
\boxtimes	is attached hereto.	
	was filed on	as Serial No
	said application hav	ving been amended on

Design Patent

I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose to the U.S. Patent and Trademark Office all information known to be material to the patentability of this application in accordance with 37 CFR § 1.56.

I hereby claim foreign priority benefits under 35 U.S.C. § 119(a-d) of any foreign application(s) for patent or inventor's certificate listed below and have also identified below any foreign application for patent or inventor's certificate having a filing date before that of any application on which priority is claimed:

any applic	ation on which priority is	claimed:	
Priority Claimed (If X'd)	Country	Serial Number	Date Filed
□ _			
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Name: Nicholas C. Nicolaides	
Mailing Address: 4 Cider Mill Court Boothwyn, PA 19061	Signature
	Date of Signature:
City/State of Actual Residence: Boothwyn, Pennsylvania	Citizenship: United States

Name: Luigi Grasso	
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Philadelphia, PA 19107	Date of Signature:
City/State of Actual Residence: Philadelphia, Pennsylvania	Citizenship:United States
Name: Philip M. Sass	
Mailing Address: 1903 Blackhawk Circle Audubon, PA 19403	Signature
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Name:	
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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In Re Application of:

Nicholas C. Nicolaides, Luigi Grasso, and

Philip M. Sass

Serial No.: Not assigned Group Art Unit: Not assigned

Filing Date: November 7, 2000 Examiner: Not assigned

For: METHODS FOR GENERATING GENETICALLY ALTERED ANTIBODY-

PRODUCING CELL LINES WITH IMPROVED ANTIBODY

CHARACTERISTICS

BOX SEQUENCE

Assistant Commissioner for Patents Washington DC 20231

STATEMENT TO SUPPORT FILING AND SUBMISSION IN ACCORDANCE WITH 37 CFR §§ 1.821 THROUGH 1.825

\bowtie	I hereby state, in accordance with the requirements of 37 C.F.R. §1.821(f), that the
	contents of the paper and computer readable copies of the Sequence Listing, submitted in accordance with 37 CFR $\S1.821(c)$ and (e) , respectively are the same.
	I hereby state that the submission filed in accordance with 37 CFR $\S1.821(g)$ does not include new matter.
	I hereby state that the submission filed in accordance with 37 CFR $\$1.821(h)$ does not include new matter or go beyond the disclosure in the international application as filed.
	I hereby state that the amendments, made in accordance with 37 CFR §1.825(a), included in the substitute sheet(s) of the Sequence Listing are supported in the application, as filed, at pages I hereby state that the substitute sheet(s) of the Sequence Listing does (do) not include new matter.
	I hereby state that the substitute copy of the computer readable form, submitted in accordance with 37 CFR §1.825(b), is the same as the amended Sequence Listing.

I hereby state that the substitute copy of the computer readable form, submitted in accordance with 37 CFR §1.825(d), contains identical data to that originally filed.

Date: 11 /7 /00

Patrick J. Farley
Registration No. 42,524

Woodcock Washburn Kurtz Mackiewicz & Norris LLP One Liberty Place - 46th Floor Philadelphia PA 19103 Telephone: (215) 568-3100 Facsimile: (215) 568-3439

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His Leu Phe Tyr Thr Leu Pro Val Arg Tyr Lys Glu Phe Gln Arg Asn \$165\$

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Leu Ser Thr Ser Gly Arg His Lys Thr Phe Ser Thr Phe Arg Ala Ser \$260\$

Phe His Ser Ala Arg Thr Ala Pro Gly Gly Val Gln Gln Thr Gly Ser 275 280 285

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Val Val Leu Asn Val Ser Val Asp Ser Glu Cys Val Asp Ile Asn Val 325 330 335

Thr Pro Asp Lys Arg Gln Ile Leu Leu Gln Glu Glu Lys Leu Leu Leu 340 345 350

Ala Val Leu Lys Thr Ser Leu Ile Gly Met Phe Asp Ser Asp Ala Asn

355 360 365

Lys Leu Asn Val Asn Gln Gln Pro Leu Leu Asp Val Glu Gly Asn Leu $370 \hspace{1cm} 375 \hspace{1cm} 380$

Val Lys Leu His Thr Ala Glu Leu Glu Lys Pro Val Pro Gly Lys Gln 385 390 395 400

Asp Asn Ser Pro Ser Leu Lys Ser Thr Ala Asp Glu Lys Arg Val Ala 405 410 415

Ser Ile Ser Arg Leu Arg Glu Ala Phe Ser Leu His Pro Thr Lys Glu 420 425 430

Ile Lys Ser Arg Gly Pro Glu Thr Ala Glu Leu Thr Arg Ser Phe Pro 435 440 445

Ser Glu Lys Arg Gly Val Leu Ser Ser Tyr Pro Ser Asp Val Ile Ser 450 455

Tyr Arg Gly Leu Arg Gly Ser Gln Asp Lys Leu Val Ser Pro Thr Asp 465 470 475

Ser Pro Gly Asp Cys Met Asp Arg Glu Lys Ile Glu Lys Asp Ser Gly \$485\$

Leu Ser Ser Thr Ser Ala Gly Ser Glu Glu Glu Phe Ser Thr Pro Glu $500 \hspace{1cm} 505 \hspace{1cm} 510 \hspace{1cm}$

Val Ala Ser Ser Phe Ser Ser Asp Tyr Asn Val Ser Ser Leu Glu Asp $515 \hspace{1.5cm} 520 \hspace{1.5cm} 525$

Pro Gly Thr Gly Gln Ser Leu Lys Pro Glu Asp His Gly Tyr Gln Cys 545 550 555 560

Lys Ala Leu Pro Leu Ala Arg Leu Ser Pro Thr Asn Ala Lys Arg Phe 565 570 575

Lys Thr Glu Glu Arg Pro Ser Asn Val Asn Ile Ser Gln Arg Leu Pro $580 \hspace{1.5cm} 585 \hspace{1.5cm} 590$

Gly Pro Gln Ser Thr Ser Ala Ala Glu Val Asp Val Ala Ile Lys Met 595 600 605

Asn Lys Arg Ile Val Leu Leu Glu Phe Ser Leu Ser Ser Leu Ala Lys

610 615 620

Arg Met Lys Gln Leu Gln His Leu Lys Ala Gln Asn Lys His Glu Leu 625 630 635 640

Ser Tyr Arg Lys Phe Arg Ala Lys Ile Cys Pro Gly Glu Asn Gln Ala \$645\$

Ala Glu Asp Glu Leu Arg Lys Glu Ile Ser Lys Ser Met Phe Ala Glu 660 665 670

Met Glu Ile Leu Gly Gln Phe Asn Leu Gly Phe Ile Val Thr Lys Leu 675 680 685

Lys Glu Asp Leu Phe Leu Val Asp Gln His Ala Ala Asp Glu Lys Tyr $690 \hspace{1.5cm} 695 \hspace{1.5cm} 700 \hspace{1.5cm}$

Asn Phe Glu Met Leu Gln Gln His Thr Val Leu Gln Ala Gln Arg Leu 705 710715715

Ile Thr Pro Gln Thr Leu Asn Leu Thr Ala Val Asn Glu Ala Val Leu 725 730 735

Ile Glu Asn Leu Glu Ile Phe Arg Lys Asn Gly Phe Asp Phe Val Ile 740 745 750

Asp Glu Asp Ala Pro Val Thr Glu Arg Ala Lys Leu Ile Ser Leu Pro $755 \hspace{1cm} 765 \hspace{1cm} 765 \hspace{1cm}$

Phe Met Leu Ser Asp Ser Pro Gly Val Met Cys Arg Pro Ser Arg Val 785 790 795 800

Arg Gln Met Phe Ala Ser Arg Ala Cys Arg Lys Ser Val Met Ile Gly 805 810 815

Thr Ala Leu Asn Ala Ser Glu Met Lys Lys Leu Ile Thr His Met Gly 820 825 830

Glu Met Asp His Pro Trp Asn Cys Pro His Gly Arg Pro Thr Met Arg 835 840 845

His Val Ala Asn Leu Asp Val Ile Ser Gln Asn 850 855 <210> 6 <211> 3056 <212> DNA <213> Mus musculus

<400> 6

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<212> PRT

<213> Homo sapiens

<400> 7

Met Glu Arg Ala Glu Ser Ser Ser Thr Glu Pro Ala Lys Ala Ile Lys $1 \hspace{1cm} 5 \hspace{1cm} 10 \hspace{1cm} 15$

Pro Ile Asp Arg Lys Ser Val His Gln Ile Cys Ser Gly Gln Val Val $20 \hspace{1.5cm} 25 \hspace{1.5cm} 30$

Leu Ser Leu Ser Thr Ala Val Lys Glu Leu Val Glu Asn Ser Leu Asp \$35\$ \$40\$ \$45\$

Ala Gly Ala Thr Asn Ile Asp Leu Lys Leu Lys Asp Tyr Gly Val Asp $50 \ \ 55 \ \ \ 60$

Glu Gly Leu Thr Leu Lys His His Thr Ser Lys Ile Glu Phe Ala $85 \hspace{1.5cm} 90 \hspace{1.5cm} 95$

Ser Leu Cys Ala Leu Ser Asp Val Thr Ile Ser Thr Cys His Ala Ser \$115\$ \$120\$ \$125\$

Ala Lys Val Gly Thr Arg Leu Met Phe Asp His Asn Gly Lys Ile Ile 130 \$135\$

Gln Lys Thr Pro Tyr Pro Arg Pro Arg Gly Thr Thr Val Ser Val Gln 145 \$150\$

Gln Leu Phe Ser Thr Leu Pro Val Arg His Lys Glu Phe Gln Arg Asn

165 170 175

Ile Lys Lys Glu Tyr Ala Lys Met Val Gln Val Leu His Ala Tyr Cys 180 185 190

Ile Ile Ser Ala Gly Ile Arg Val Ser Cys Thr Asn Gln Leu Gly Gln $195 \hspace{1cm} 200 \hspace{1cm} 205 \hspace{1cm}$

Gly Lys Arg Gln Pro Val Val Cys Thr Gly Gly Ser Pro Ser Ile Lys 210 215 220

Glu Asn Ile Gly Ser Val Phe Gly Gln Lys Gln Leu Gln Ser Leu Ile 225 230 235 240

Pro Phe Val Gln Leu Pro Pro Ser Asp Ser Val Cys Glu Glu Tyr Gly \$245\$ \$250\$

Ile Ser Gln Cys Thr His Gly Val Gly Arg Ser Ser Thr Asp Arg Gln \$275\$

Phe Phe Phe Ile Asn Arg Arg Pro Cys Asp Pro Ala Lys Val Cys Arg 290 295 300

Leu Val Asn Glu Val Tyr His Met Tyr Asn Arg His Gln Tyr Pro Phe 305 310 315

Val Val Leu Asn Ile Ser Val Asp Ser Glu Cys Val Asp Ile Asn Val 325 330 335

Thr Pro Asp Lys Arg Gln Ile Leu Leu Gln Glu Glu Lys Leu Leu Leu 340 345 350

Ala Val Leu Lys Thr Ser Leu Ile Gly Met Phe Asp Ser Asp Val Asn 355 360 365

Lys Leu Asn Val Ser Gln Gln Pro Leu Leu Asp Val Glu Gly Asn Leu 370 375 380

Ile Lys Met His Ala Ala Asp Leu Glu Lys Pro Met Val Glu Lys Gln 385 \$390\$ 395 400

Asp Gln Ser Pro Ser Leu Arg Thr Gly Glu Glu Lys Lys Asp Val Ser 405 410 415

Ile Ser Arg Leu Arg Glu Ala Phe Ser Leu Arg His Thr Thr Glu Asn

Lys Pro His Ser Pro Lys Thr Pro Glu Pro Arg Arg Ser Pro Leu Gly 435 440 445

Gln Lys Arg Gly Met Leu Ser Ser Ser Thr Ser Gly Ala Ile Ser Asp $450 \ \ 455 \ \ 460$

Lys Gly Val Leu Arg Pro Gln Lys Glu Ala Val Ser Ser Ser His Gly 465 470 475 480

Pro Ser Asp Pro Thr Asp Arg Ala Glu Val Glu Lys Asp Ser Gly His \$485\$

Gly Ser Thr Ser Val Asp Ser Glu Gly Phe Ser Ile Pro Asp Thr Gly $500 \hspace{1.5cm} 505 \hspace{1.5cm} 510 \hspace{1.5cm}$

Ser His Cys Ser Ser Glu Tyr Ala Ala Ser Ser Pro Gly Asp Arg Gly 515 520 525

Ser Gln Glu His Val Asp Ser Gln Glu Lys Ala Pro Glu Thr Asp Asp 530 540

Ser Phe Ser Asp Val Asp Cys His Ser Asn Gln Glu Asp Thr Gly Cys 545 550 555 560

Lys Phe Arg Val Leu Pro Gln Pro Thr Asn Leu Ala Thr Pro Asn Thr \$565\$ \$570\$ \$575\$

Lys Arg Phe Lys Lys Glu Glu Ile Leu Ser Ser Ser Asp Ile Cys Gln 580 585 590

Lys Leu Val Asn Thr Gln Asp Met Ser Ala Ser Gln Val Asp Val Ala $595 \hspace{1.5cm} 600 \hspace{1.5cm} 605 \hspace{1.5cm}$

Val Lys Ile Asn Lys Lys Val Val Pro Leu Asp Phe Ser Met Ser Ser 610 615 620

Leu Ala Lys Arg Ile Lys Gln Leu His His Glu Ala Gln Gln Ser Glu 625 630 635 640

Gly Glu Gln Asn Tyr Arg Lys Phe Arg Ala Lys Ile Cys Pro Gly Glu $645 \hspace{1cm} 650 \hspace{1cm} 655$

Asn Gln Ala Ala Glu Asp Glu Leu Arg Lys Glu Ile Ser Lys Thr Met \$660\$

Phe Ala Glu Met Glu Ile Ile Gly Gln Phe Asn Leu Gly Phe Ile Ile

Thr Lys Leu Asn Glu Asp Ile Phe Ile Val Asp Gln His Ala Thr Asp 690 695 700

Glu Lys Tyr Asn Phe Glu Met Leu Gln Gln His Thr Val Leu Gln Gly 705 $710715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715715$

Gln Arg Leu Ile Ala Pro Gln Thr Leu Asn Leu Thr Ala Val Asn Glu 725 730 735

Ala Val Leu Ile Glu Asn Leu Glu Ile Phe Arg Lys Asn Gly Phe Asp 740 745 750

Phe Val Ile Asp Glu Asn Ala Pro Val Thr Glu Arg Ala Lys Leu Ile $755 \hspace{1.5cm} 760 \hspace{1.5cm} 765 \hspace{1.5cm}$

Ser Leu Pro Thr Ser Lys Asn Trp Thr Phe Gly Pro Gln Asp Val Asp 770 775 780

Glu Leu Ile Phe Met Leu Ser Asp Ser Pro Gly Val Met Cys Arg Pro 785 790 795

Ser Arg Val Lys Gln Met Phe Ala Ser Arg Ala Cys Arg Lys Ser Val\$805\$ \$810\$ \$10

Met Ile Gly Thr Ala Leu Asn Thr Ser Glu Met Lys Lys Leu Ile Thr 820 825 830

His Met Gly Glu Met Asp His Pro Trp Asn Cys Pro His Gly Arg Pro 835 840 845

Thr Met Arg His Ile Ala Asn Leu Gly Val Ile Ser Gln Asn 850 860

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tcactttgtg cactgagcga tgtcaccatt tctacctgcc acgcatcggc gaaggttgga 420 actegactga tgtttgatca caatgggaaa attatecaga aaacccccta cccccgcccc 480 agagggacca cagtcagcgt gcagcagtta ttttccacac tacctgtgcg ccataaggaa 540 tttcaaagga atattaagaa ggagtatgcc aaaatggtcc aggtcttaca tgcatactgt 600 atcatttcag caggcatccg tgtaagttgc accaatcagc ttggacaagg aaaacgacag 660 cctgtggtat gcacaggtgg aagccccage ataaaggaaa atatcggctc tgtgtttggg 720 cagaaqcaqt tqcaaaqcct cattcctttt gttcagctgc cccctagtga ctccgtgtgt 780 gaagagtacg gtttgagctg ttcggatgct ctgcataatc ttttttacat ctcaggtttc 840 atttcacaat gcacgcatgg agttggaagg agttcaacag acagacagtt tttctttatc 900 aaccggcggc cttgtgaccc agcaaaggtc tgcagactcg tgaatgaggt ctaccacatg 960 tataatcgac accagtatcc atttgttgtt cttaacattt ctgttgattc agaatgcgtt 1020 gatatcaatg ttactccaga taaaaggcaa attttgctac aagaggaaaa gcttttgttg 1080 geagttttaa agacetettt gataggaatg tttgatagtg atgteaacaa getaaatgte 1140 agtcagcagc cactgctgga tgttgaaggt aacttaataa aaatgcatgc agcggatttg 1200 gaaaagccca tggtagaaaa gcaggatcaa tccccttcat taaggactgg agaagaaaaa 1260 aaagacgtgt ccatttccag actgcgagag gccttttctc ttcgtcacac aacagagaac 1320 aagcctcaca gcccaaagac tccagaacca agaaggagcc ctctaggaca gaaaaggggt 1380 atgctqtctt ctaqcacttc aqqtqccatc tctqacaaaq qcqtcctqaq acctcaqaaa 1440 gaggcagtga gttccagtca cggacccagt gaccctacgg acagagcgga ggtggagaag 1500 gactegggge acggeageac tteegtggat tetgaggggt teageatece agacaeggge 1560 agtcactgca gcaqcqaqta tgcggccaqc tccccaqqqq acaqqqqctc qcaqqaacat 1620 gtggactete aggagaaage geetgaaact gaegaetett titeagatgt ggaetgeeat 1680 tcaaaccagg aagataccgg atgtaaattt cgagttttgc ctcagccaac taatctcgca 1740 accccaaaca caaagcgttt taaaaaagaa gaaattettt ecagttetga catttgteaa 1800 aagttagtaa atactcagga catgtcagcc tctcaggttg atgtagctgt gaaaattaat 1860 aaqaaaqttq tqcccctgga cttttctatg aqttctttag ctaaacqaat aaaqcaqtta 1920 catcatgaag cacagcaaag tgaaggggaa cagaattaca ggaagttag ggcaaagatt 1980 tgtcctggag aaaatcaagc agccgaagat gaactaagaa aagagataag taaaacgatg 2040 tttqcaqaaa tqqaaatcat tqqtcaqttt aacctqqqat ttataataac caaactqaat 2100 gaggatatet teatagtgga ceageatgee acggacgaga agtataaett egagatgetg 2160 cagcagcaca ccgtgctcca ggggcagagg ctcatagcac ctcagactct caacttaact 2220 gctgttaatg aagctgttct gatagaaaat ctggaaatat ttagaaagaa tggctttgat 2280 tttgttatcg atgaaaatgc tccagtcact gaaagggcta aactgatttc cttgccaact 2340 agtaaaaact ggaccttcgg accccaggac gtcgatgaac tgatcttcat gctgagcgac 2400 agccetgggg teatgtgeeg geetteegga gteaageaga tgtttgeete eagageetge 2460 cggaaqtcqq tqatqattqq qactgctctt aacacaaqcq aqatqaaqaa actqatcacc 2520 cacatggggg agatggacca ccctggaac tgtccccatg gaaggccaac catgagacac 2580 atogocaaco tgggtgtcat ttotcagaac tgaccgtagt cactgtatgg aataattggt 2640 tttatcgcag atttttatgt tttgaaagac agagtcttca ctaacctttt ttgttttaaa 2700 atgaaacctg ctacttaaaa aaaatacaca tcacacccat ttaaaagtga tcttgagaac 2760

2771

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<212> PRT

<213> Homo sapiens

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Ile	Ile	Thr	Ser 20	Val	Val	Ser	Val	Val 25	Lys	Glu	Leu	Ile	Glu 30	Asn	Ser
Leu	Asp	Ala 35	Gly	Ala	Thr	Ser	Val 40	Asp	Val	Lys	Leu	Glu 45	Asn	Tyr	Gly
Phe	Asp 50	Lys	Ile	Glu	Val	Arg 55	Asp	Asn	Gly	Glu	Gly 60	Ile	Lys	Ala	Val
Asp 65	Ala	Pro	Val	Met	Ala 70	Met	Lys	Tyr	Tyr	Thr 75	Ser	Lys	Ile	Asn	Ser 80
His	Glu	Asp	Leu	Glu 85	Asn	Leu	Thr	Thr	Tyr 90	Gly	Phe	Arg	Gly	Glu 95	Ala
Leu	Gly	Ser	Ile 100	Cys	Cys	Ile	Ala	Glu 105	Val	Leu	Ile	Thr	Thr 110	Arg	Thr
Ala	Ala	Asp		Phe					Val		_	Gly		Gly	His

115 120 125

Ile Leu Ser Gln Lys Pro Ser His Leu Gly Gln Gly Thr Thr Val Thr 130 135 140

Ala Leu Arg Leu Phe Lys Asn Leu Pro Val Arg Lys Gln Phe Tyr Ser 145 \$150\$

Thr Ala Lys Lys Cys Lys Asp Glu Ile Lys Lys Ile Gln Asp Leu Leu $165 \hspace{1.5cm} 170 \hspace{1.5cm} 175$

Met Ser Phe Gly Ile Leu Lys Pro Asp Leu Arg Ile Val Phe Val His 180 $$185\$

Asn Lys Ala Val Ile Trp Gln Lys Ser Arg Val Ser Asp His Lys Met 195 \$200\$

Ala Leu Met Ser Val Leu Gly Thr Ala Val Met Asn Asn Met Glu Ser 210 215 220

Phe Gln Tyr His Ser Glu Glu Ser Gln Ile Tyr Leu Ser Gly Phe Leu 225 230 235 240

Pro Lys Cys Asp Ala Asp His Ser Phe Thr Ser Leu Ser Thr Pro Glu \$245\$

Arg	Ser	Phe	Ile 260	Phe	Ile	Asn	Ser	Arg 265	Pro	Val	His	Gln	Lys 270	Asp	Ile
Leu	Lys	Leu 275	Ile	Arg	His	His	Tyr 280	Asn	Leu	Lys	Cys	Leu 285	Lys	Glu	Ser
Thr	Arg 290	Leu	Tyr	Pro	Val	Phe 295	Phe	Leu	Lys	Ile	Asp 300	Val	Pro	Thr	Ala
Asp 305	Val	Asp	Val	Asn	Leu 310	Thr	Pro	Asp	Lys	Ser 315	Gln	Val	Leu	Leu	Glr 320
Asn	Lys	Glu	Ser	Val 325	Leu	Ile	Ala	Leu	G1u 330	Asn	Leu	Met	Thr	Thr 335	Cys
			Leu 340					345					350		
		355	Ala				360					365			
	370		Lys			375					380				
385			Ile		390					395					400
			Asp	405					410				_	415	
			His 420					425					430		
		435	Phe		-		440					445			
	450		Glu	-		455		-			460			-	
465			Glu		470		-	-		475	-			-	480
			Glu Arg	485					490					495	
JIU	ı.p	Set	500	эту	Hoil	116	±eu.	505	ASII	ner.	Val	этУ	510	ASII	тте

Glu Pro Val Lys Ile Leu Val Pro Glu Lys Ser Leu Pro Cys Lys Val Ser Asn Asn Asn Tyr Pro Ile Pro Glu Gln Met Asn Leu Asn Glu Asp Ser Cys Asn Lys Lys Ser Asn Val Ile Asp Asn Lys Ser Gly Lys Val Thr Ala Tyr Asp Leu Leu Ser Asn Arg Val Ile Lys Lys Pro Met Ser Ala Ser Ala Leu Phe Val Gln Asp His Arg Pro Gln Phe Leu Ile Glu Asn Pro Lys Thr Ser Leu Glu Asp Ala Thr Leu Gln Ile Glu Glu Leu Trp Lys Thr Leu Ser Glu Glu Glu Lys Leu Lys Tyr Glu Glu Lys Ala Thr Lys Asp Leu Glu Arg Tyr Asn Ser Gln Met Lys Arg Ala Ile Glu Gln Glu Ser Gln Met Ser Leu Lys Asp Gly Arg Lys Lys Ile Lys Pro Thr Ser Ala Trp Asn Leu Ala Gln Lys His Lys Leu Lys Thr Ser Leu Ser Asn Gln Pro Lys Leu Asp Glu Leu Leu Gln Ser Gln Ile Glu Lys Arg Arg Ser Gln Asn Ile Lys Met Val Gln Ile Pro Phe Ser Met Lys Asn Leu Lys Ile Asn Phe Lys Lys Gln Asn Lys Val Asp Leu Glu Glu Lys Asp Glu Pro Cys Leu Ile His Asn Leu Arg Phe Pro Asp Ala Trp Leu Met Thr Ser Lys Thr Glu Val Met Leu Leu Asn Pro Tyr Arg Val

Glu Glu Ala Leu Leu Phe Lys Arg Leu Leu Glu Asn His Lys Leu Pro
755 760 765

Ala Glu Pro Leu Glu Lys Pro Ile Met Leu Thr Glu Ser Leu Phe Asn 770 780

Gly Ser His Tyr Leu Asp Val Leu Tyr Lys Met Thr Ala Asp Asp Gln 785 790 795

Arg Tyr Ser Gly Ser Thr Tyr Leu Ser Asp Pro Arg Leu Thr Ala Asn 805 810 815

Gly Phe Lys Ile Lys Leu Ile Pro Gly Val Ser Ile Thr Glu Asn Tyr 820 825 830

Leu Glu Ile Glu Gly Met Ala Asn Cys Leu Pro Phe Tyr Gly Val Ala 835 840 845

Asp Leu Lys Glu Ile Leu Asn Ala Ile Leu Asn Arg Asn Ala Lys Glu 850 855 860

Val Tyr Glu Cys Arg Pro Arg Lys Val Ile Ser Tyr Leu Glu Gly Glu 865 870 875 880

Ala Val Arg Leu Ser Arg Gln Leu Pro Met Tyr Leu Ser Lys Glu Asp 885 890 895

Ile Gln Asp Ile Ile Tyr Arg Met Lys His Gln Phe Gly Asn Glu Ile $900 \hspace{1.5cm} 905 \hspace{1.5cm} 910$

Lys Glu Cys Val His Gly Arg Pro Phe Phe His His Leu Thr Tyr Leu \$915\$ \$920\$ \$925

Pro Glu Thr Thr 930

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<213> Homo sapiens

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<213> Homo sapiens

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Thr Val Arg Leu Phe Asp Arg Gly Asp Phe Tyr Thr Ala His Gly Glu $35 \hspace{1cm} 40 \hspace{1cm} 45$

Lys Tyr Met Gly Pro Ala Gly Ala Lys Asn Leu Gln Ser Val Val Leu 65 70 75 80

Ser Lys Met Asn Phe Glu Ser Phe Val Lys Asp Leu Leu Leu Val Arg 85 90 95

Gln Tyr Arg Val Glu Val Tyr Lys Asn Arg Ala Gly Asn Lys Ala Ser $100 \\ 105 \\ 110$

Lys Glu Asn Asp Trp Tyr Leu Ala Tyr Lys Ala Ser Pro Gly Asn Leu \$115\$ \$120\$

Ser Gln Phe Glu Asp Ile Leu Phe Gly Asn Asn Asp Met Ser Ala Ser 130 135

Ile Gly Val Val Gly Val Lys Met Ser Ala Val Asp Gly Gln Arg Gln 145 \$150\$

Val Gly Val Gly Tyr Val Asp Ser Ile Gln Arg Lys Leu Gly Leu Cys $165 \hspace{1.5cm} 170 \hspace{1.5cm} 175$

Glu Phe Pro Asp Asp Asp Gln Phe Ser Asn Leu Glu Ala Leu Leu Ile 180 $$185\$

Gln Ile Gly Pro Lys Glu Cys Val Leu Pro Gly Gly Glu Thr Ala Gly 195 \$200\$

Asp Met Gly Lys Leu Arg Gln Ile Ile Gln Arg Gly Gly Ile Leu Ile 210 215 220

Thr Glu Arg Lys Lys Ala Asp Phe Ser Thr Lys Asp Ile Tyr Gln Asp

Leu Asn Arg Leu Leu Lys Gly Lys Lys Gly Glu Gln Met Asn Ser Ala 245 250 255

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Phe Glu Leu Thr Thr Phe Asp Phe Ser Gln Tyr Met Lys Leu Asp Ile 290 295 300

Ala Ala Val Arg Ala Leu Asn Leu Phe Gln Gly Ser Val Glu Asp Thr 305 310 315 320

Thr Gly Ser Gln Ser Leu Ala Ala Leu Leu Asn Lys Cys Lys Thr Pro 325 330 335

Gln Gly Gln Arg Leu Val Asn Gln Trp Ile Lys Gln Pro Leu Met Asp \$340\$ \$345\$ \$350

Lys Asn Arg Ile Glu Glu Arg Leu Asn Leu Val Glu Ala Phe Val Glu 355 \$360\$

Pro Asp Leu Asn Arg Leu Ala Lys Lys Phe Gln Arg Gln Ala Ala Asn 385 390 395 400

Leu Gln Asp Cys Tyr Arg Leu Tyr Gln Gly Ile Asn Gln Leu Pro Asn 405 410 415

Val Ile Gln Ala Leu Glu Lys His Glu Gly Lys His Gln Lys Leu Leu \$420\$ \$425\$

Leu Ala Val Phe Val Thr Pro Leu Thr Asp Leu Arg Ser Asp Phe Ser \$435\$

Lys Phe Gln Glu Met Ile Glu Thr Thr Leu Asp Met Asp Gln Val Glu $450 \,$ $\,$ $455 \,$ $\,$ $460 \,$

Asn His Glu Phe Leu Val Lys Pro Ser Phe Asp Pro Asn Leu Ser Glu 465 470 475 480

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485 490 495

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500 505 510

Leu Asp Ser Ser Ala Gln Phe Gly Tyr Tyr Phe Arg Val Thr Cys Lys 515 520 525

Glu Glu Lys Val Leu Arg Asn Asn Lys Asn Phe Ser Thr Val Asp Ile 530 535 540

Gln Lys Asn Gly Val Lys Phe Thr Asn Ser Lys Leu Thr Ser Leu Asn 545 550 555 560

Glu Glu Tyr Thr Lys Asn Lys Thr Glu Tyr Glu Glu Ala Gln Asp Ala 565 570 575

Ile Val Lys Glu Ile Val Asn Ile Ser Ser Gly Tyr Val Glu Pro Met $580 \hspace{1.5cm} 585 \hspace{1.5cm} 590 \hspace{1.5cm}$

Gln Thr Leu Asn Asp Val Leu Ala Gln Leu Asp Ala Val Val Ser Phe $595 \hspace{1.5cm} 600 \hspace{1.5cm} 605$

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Cys Val Glu Val Gln Asp Glu Ile Ala Phe Ile Pro Asn Asp Val Tyr \$645\$

Gly Gly Lys Ser Thr Tyr Ile Arg Gln Thr Gly Val Ile Val Leu Met 675 680 685

Ala Gln Ile Gly Cys Phe Val Pro Cys Glu Ser Ala Glu Val Ser Ile 690 695 700

Val Asp Cys Ile Leu Ala Arg Val Gly Ala Gly Asp Ser Gln Leu Lys 705 710 715 720

Gly Val Ser Thr Phe Met Ala Glu Met Leu Glu Thr Ala Ser Ile Leu 725 730 735

Arg Ser Ala Thr Lys Asp Ser Leu Ile Ile Ile Asp Glu Leu Gly Arg

750

Gly Thr Ser Thr Tyr Asp Gly Phe Gly Leu Ala Trp Ala Ile Ser Glu 755 760 765

Tyr Ile Ala Thr Lys Ile Gly Ala Phe Cys Met Phe Ala Thr His Phe 770 775 780

His Glu Leu Thr Ala Leu Ala Asn Gln Ile Pro Thr Val Asn Asn Leu 785 790 795 800

His Val Thr Ala Leu Thr Thr Glu Glu Thr Leu Thr Met Leu Tyr Gln 805 810 815

Val Lys Lys Gly Val Cys Asp Gln Ser Phe Gly Ile His Val Ala Glu 820 825 830

Leu Ala Asn Phe Pro Lys His Val Ile Glu Cys Ala Lys Gln Lys Ala $835 \\ 840 \\ 845$

Leu Glu Leu Glu Glu Phe Gln Tyr Ile Gly Glu Ser Gln Gly Tyr Asp 850

Ile Met Glu Pro Ala Ala Lys Lys Cys Tyr Leu Glu Arg Glu Gln Gly 865 870 875

Glu Lys Ile Ile Gln Glu Phe Leu Ser Lys Val Lys Gln Met Pro Phe \$885\$

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<212> DNA

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<212> PRT

<213> Homo sapiens

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Lys Glu Met Ile Glu Asn Cys Leu Asp Ala Lys Ser Thr Ser Ile Glu \$35\$ \$40\$ \$45\$

Val Ile Val Lys Glu Gly Gly Leu Lys Leu Ile Gln Ile Gln Asp Asn 50 55 60

Gly Thr Gly Ile Arg Lys Glu Asp Leu Asp Ile Val Cys Glu Arg Phe 65 70 75 80

Thr Thr Ser Lys Leu Gln Ser Phe Glu Asp Leu Ala Ser Ile Ser Thr 85 90 95

Val Thr Ile Thr Thr Lys Thr Ala Asp Gly Lys Cys Ala Tyr Arg Ala 115 120 125

Ser Tyr Ser Asp Gly Lys Leu Lys Ala Pro Pro Lys Pro Cys Ala Gly 130 $$135\$

Asn Gln Gly Thr Gln Ile Thr Val Glu Asp Leu Phe Tyr Asn Ile Ala 145 \$150\$

Thr Arg Arg Lys Ala Leu Lys Asn Pro Ser Glu Glu Tyr Gly Lys Ile 165 170 175

Leu Glu Val Val Gly Arg Tyr Ser Val His Asn Ala Gly Ile Ser Phe 180 185 190

Ser Val Lys Lys Gln Gly Glu Thr Val Ala Asp Val Arg Thr Leu Pro \$195\$

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Ser	Lys	Pro	Leu	Ser 405	Ser	Gln	Pro	Gln	Ala 410	Ile	Val	Thr	Glu	Asp 415	Lys
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455

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Pro 705	Gly	Ser	Ile	Pro	Asn 710	Ser	Trp	Lys	Trp	Thr 715	Val	Glu	His	Ile	Val 720

Tyr Lys Ala Leu Arg Ser His Ile Leu Pro Pro Lys His Phe Thr Glu 725 730 735

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<210> 15

<211> 133

<212> PRT

<213> Homo sapiens

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Leu Ser Leu Ser Thr Ala Val Lys Glu Leu Val Glu Asn Ser Leu Asp \$35\$ \$40\$ \$45\$

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Glu Gly Leu Thr Leu Lys His His Thr Ser Lys Ile Gln Glu Phe Ala \$85\$ 90 95

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